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CHALCONE SYNTHASE DIHYDROFLAVONOL 4-REDUCTASE AND LEUCOANTHOCYANIDINE REDUCTASE FROM CLOVER, MEDIC RYEGRASS OR FESCUE

(57) Abstract: The present invention relates to nucleic acid fragments encoding amino acid sequences for flavonoid biosynthetic enzymes in plants, and the use thereof for the modification of, for example, flavonoid biosynthesis in plants, and more specifically the modification of the content of condensed tannins. In particularly preferred embodiments, the invention relates to the combinatorial expression of chalcone synthase (CHS) and/or dihydroflavonol 4-reductase (BAN) and/or leucoanthocyanidine reductase (LAR) in plants to modify, for example, flavonoid biosynthesis or more specifically the content of condensed tannins.

Chalcone Synthase, Dihydroflavonol 4-reductase and Leucoanthocyanidine reductase from Clover, Medic, Ryegrass or Fescue.

The present invention relates to nucleic acid fragments encoding amino acid sequences for flavonoid biosynthetic enzyme polypeptides in plants, and the use thereof for the modification of, for example, flavonoid biosynthesis in plants, and more specifically the modification of the content of condensed tannins. In particularly preferred embodiments, the invention relates to the combinatorial expression of chalcone synthase (CHS) and/or dihydroflavonol 4-reductase (BAN) and/or leucoanthocyanidine reductase (LAR) in plants to modify, for example, flavonoid biosynthesis or more specifically the content of condensed tannins.

Flavonoids constitute a relatively diverse family of aromatic molecules that are derived from phenylalanine and malonyl-coenzyme A (CoA, via the fatty acid pathway). These compounds include six major subgroups that are found in most higher plants: the chalcones, flavones, flavonols, flavandiols, anthocyanins and condensed tannins (or proanthocyanidins). A seventh group, the aurones, is widespread, but not ubiquitous.

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Some plant species also synthesize specialised forms of flavonoids, such as the isoflavonoids that are found in legumes and a small number of non-legume plants. Similarly, sorghum, maize and gloxinia are among the few species known to synthesize 3-deoxyanthocyanins (or phlobaphenes in the polymerised form). The stilbenes, which are closely related to flavonoids, are synthesised by another group of unrelated species that includes grape, peanut and pine.

Besides providing pigmentation to flowers, fruits, seeds, and leaves, flavonoids also have key roles in signalling between plants and microbes, in male fertility of some species, in defence as antimicrobial agents and feeding deterrents, and in UV protection.

Flavonoids also have significant activities when ingested by animals, and there is great interest in their potential health benefits, particularly for compounds such as isoflavonoids, which have been linked to anticancer benefits, and stilbenes that are believed to contribute to reduced heart disease. Condensed tannins which are plant polyphenols with protein-precipitating and antioxidant properties are involved in protein binding, metal chelation, anti-oxidation, and UV-light absorption. As a result condensed tannins inhibit viruses, microorganisms, insects, fungal pathogens, and monogastric digestion. Moderate amounts of

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tannins improve forage quality by disrupting protein foam and conferring protection from rumen pasture bloat. Bloat is a digestive disorder that occurs on some highly nutritious forage legumes such as alfalfa (*Medicago sativa*) and white clover (*Trifolium repens*). Moderate amounts of tannin can also reduce digestion rates in the rumen and can reduce parasitic load sufficiently to increase the titre of amino acids and small peptides in the small intestine without compromising total digestion.

The major branch pathways of flavonoid biosynthesis start with general phenylpropanoid metabolism and lead to the nine major subgroups: the colourless chalcones, aurones, isoflavonoids, flavones, flavonols, flavandiols, anthocyanins, condensed tannins, and phlobaphene pigments. The enzyme phenylalanine ammonia-lyase (PAL) of the general phenylpropanoid pathway will lead to the production of cinnamic acid. Cinnamate-4-hydroxylase (C4H) will produce p-coumaric acid which will be converted through the action of 4-coumaroyl:CoA-ligase (4CL) to the production of 4-coumaroyl-CoA and malonyl-CoA. The first committed step channelling carbon into the flavonoid biosynthesis pathway is catalysed by chalcone synthase (CHS), which uses malonyl CoA and 4-coumaryl CoA as substrates.

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The Arabidopsis BANYULS gene encodes a dihydroflavonol 4-reductase-like protein (BAN) that may be an anthocyanine reductase (ACR). The reaction catalysed by BAN is considered to be one possible branching point from the general flavonoid pathway to the condensed tannin biosynthesis.

An alternative pathway to condensed tannins is via leucoanthocyanidine reductase (LAR). LAR utilises the same substrate as the ACR (BAN) but produces a 2,3-trans isomer as compared to the 2,3-cis isomer produced by ACR.

While nucleic acid sequences encoding the key enzymes in the condensed tannins biosynthetic pathway CHS, BAN and LAR have been isolated for certain species of plants, there remains a need for materials useful in modifying flavonoid biosynthesis and more specifically in modifying condensed tannin biosynthesis and therewith in modifying forage quality, for example by disrupting protein foam and conferring protection from rumen pasture bloat, particularly in forage legumes and grasses, including alfalfa, medics, clovers, ryegrasses and fescues, and for methods for their use.

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It is an object of the present invention to overcome, or at least alleviate, one or more of the difficulties or deficiencies associated with the prior art.

In one aspect, the present invention provides substantially purified or isolated nucleic acids or nucleic acid fragments encoding key polypeptide enzymes in the condensed tannins biosynthetic pathway CHS, BAN and LAR, or functionally active fragments or variants of these enzymes, from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species.

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The present invention also provides substantially purified or isolated nucleic acids or nucleic acid fragments encoding amino acid sequences for a class of polypeptides which are related to CHS, BAN and LAR or functionally active fragments or variants of CHS, BAN or LAR. Such polypeptides are referred to herein as CHS-like, BAN-like and LAR-like, respectively, and includes polypeptides having similar functional activity.

The individual or simultaneous enhancement or otherwise manipulation of CHS, BAN and LAR or like gene activities in plants may enhance or otherwise alter flavonoid biosynthesis; may enhance or otherwise alter the plant capacity for protein binding, metal chelation, anti-oxidation, and UV-light absorption; may enhance or reduce or otherwise alter plant pigment production; and may enhance or otherwise alter the amount of condensed tannins contained within forage legumes and grasses, including alfalfa, medics, clovers, ryegrasses and fescues and therewith the capacity to reduce bloating by disrupting protein foam.

Methods for the manipulation of CHS, BAN and LAR or like gene activities in plants, including legumes such as clovers (*Trifolium* species), lucerne (*Medicago sativa*) and grass species such as ryegrasses (*Lolium* species) and fescues (*Festuca* species) may facilitate the production of, for example, forage legumes and forage grasses and other crops with enhanced tolerance to biotic stresses such as viruses, microorganisms, insects and fungal pathogens; altered pigmentation in flowers; forage legumes with enhanced herbage quality and bloat-safety.

The clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species may be of any suitable type, including white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium*)

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subterraneum), alfalfa (Medicago sativa), Italian or annual ryegrass (Lolium multiflorum), perennial ryegrass (Lolium perenne), tall fescue (Festuca arundinacea), meadow fescue (Festuca pratensis) and red fescue (Festuca rubra). Preferably the species is a clover or a ryegrass, more preferably white clover (T. repens) or perennial ryegrass (L. perenne). White clover (Trifolium repens L.) and perennial ryegrass (Lolium perenne L.) are key pasture legumes and grasses, respectively, in temperate climates throughout the world. Perennial ryegrass is also an important turf grass.

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The nucleic acid or nucleic acid fragment may be of any suitable type and includes DNA (such as cDNA or genomic DNA) and RNA (such as mRNA) that is single- or double-stranded, optionally containing synthetic, non-natural or altered nucleotide bases, and combinations thereof. The RNA is readily obtainable, for example, by transcription of a DNA sequence according to the present invention, to produce an RNA corresponding to the DNA sequence. The RNA may be synthesised, *in vivo* or *in vitro* or by chemical synthesis to produce a sequence corresponding to a DNA sequence by methods well known in the art. In this specification, where the degree of sequence similarity between an RNA and DNA is such that the strand of the DNA could encode the RNA, then the RNA is said to "correspond" to that DNA.

In a preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a CHS or CHS-like protein includes the nucleotide sequences shown in Figures 2, 6, 10 and 14 hereto (Sequence ID Nos. 1, 3, 5 and 7, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c), and (d).

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a BAN or BAN-like protein includes the nucleotide sequence shown in Figure 18 hereto (Sequence ID No. 9); (b) complements of the sequence recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally

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active fragments and variants of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c), and (d).

In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated nucleic acid or nucleic acid fragment encoding a LAR or LAR-like protein includes the nucleotide sequence shown in Figures 22, 26 and 30 hereto (Sequence ID Nos. 11, 13 and 15 respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c), and (d).

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The term "isolated" means that the material is removed from its original environment (e.g. the natural environment if it is naturally occurring). For example, a naturally occurring nucleic acid or polypeptide present in a living plant is not isolated, but the same nucleic acid or polypeptide separated from some or all of the coexisting materials in the natural system, is isolated. Such nucleic acids could be part of a vector and/or such nucleic acids could be part of a composition, and still be isolated in that such a vector or composition is not part of its natural environment. An isolated polypeptide could be part of a composition and still be isolated in that such a composition is not part of its natural environment.

The term "purified" means that the nucleic acid or polypeptide is substantially free of other nucleic acids or polypeptides.

By "functionally active" in respect of a nucleic acid it is meant that the fragment or variant (such as an analogue, derivative or mutant) is capable of modifying flavonoid biosynthesis in a plant. Such variants include naturally occurring allelic variants and non-naturally occurring variants. Additions, deletions, substitutions and derivatizations of one or more of the nucleotides are contemplated so long as the modifications do not result in loss of functional activity of the fragment or variant. Preferably the functionally active fragment or variant has at least approximately 80% identity to the relevant part of the above mentioned sequence, more preferably at least approximately 90% identity, most preferably at least approximately 95% identity. Such functionally active variants and fragments include, for example, those having nucleic acid changes which result in conservative amino acid substitutions of one or more residues in the

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corresponding amino acid sequence. Preferably the fragment has a size of at least 30 nucleotides, more preferably at least 45 nucleotides, most preferably at least 60 nucleotides.

By "functionally active" in respect of a polypeptide is meant that the fragment or variant has one or more of the biological properties or functions of the polypeptides CHS, CHS-like, BAN, BAN-like, LAR and LAR-like, respectively. Additions, deletions, substitutions and derivatizations of one or more of the amino acids are contemplated so long as the modifications do not result in loss of functional activity of the fragment or variant. Preferably the functionally active fragment or variant has at least approximately 60% identity to the relevant part of the above mentioned sequence, more preferably at least approximately 80% identity, most preferably at least approximately 90% identity. Such functionally active variants and fragments include, for example, those having conservative amino acid substitutions of one or more residues in the corresponding amino acid sequence. Preferably the fragment has a size of at least 10 amino acids, more preferably at least 15 amino acids, most preferably at least 20 amino acids.

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The term "construct" as used herein refers to an artificially assembled or isolated nucleic acid molecule which includes the gene of interest. In general a construct may include the gene or genes of interest, a marker gene which in some cases can also be the gene of interest and appropriate regulatory sequences. It should be appreciated that the inclusion of regulatory sequences in a construct is optional, for example, such sequences may not be required in situations where the regulatory sequences of a host cell are to be used. The term construct includes vectors but should not be seen as being limited thereto.

The term "vector" as used herein encompasses both cloning and expression vectors. Vectors are often recombinant molecules containing nucleic acid molecules from several sources.

By "operatively linked" is meant that said regulatory element(s) is capable of causing expression of said nucleic acid(s) or nucleic acid fragment(s) in a plant cell and said terminator(s) is capable of terminating expression of said nucleic acid(s) or nucleic acid fragment(s) in a plant cell. Preferably, said regulatory element(s) is upstream of said nucleic acid(s) or nucleic acid fragment(s) and said terminator(s) is downstream of said nucleic acid(s) or nucleic acid fragment(s). In

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a particularly preferred embodiment, each nucleic acid or nucleic acid fragment has one or more upstream promoters and one or more downstream terminators, although expression of more than one nucleic acid or nucleic acid fragment from an upstream regulatory element(s) or termination of more than one nucleic acid or nucleic acid fragment from a downstream terminator(s) is not precluded.

By "an effective amount" it is meant an amount sufficient to result in an identifiable phenotypic trait in said plant, or a plant, plant seed or other plant part derived therefrom. Such amounts can be readily determined by an appropriately skilled person, taking into account the type of plant, the route of administration and other relevant factors. Such a person will readily be able to determine a suitable amount and method of administration. See, for example, Sambrook *et al.*, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, the entire disclosure of which is incorporated herein by reference.

It will also be understood that the term "comprises" (or its grammatical variants) as used in this specification is equivalent to the term "includes" and should not be taken as excluding the presence of other elements or features.

Genes encoding other CHS or CHS-like, BAN or BAN-like and LAR or LAR-like proteins, either as cDNAs or genomic DNAs, may be isolated directly by using all or a portion of the nucleic acids or nucleic acid fragments of the present invention as hybridisation probes to screen libraries from the desired plant employing the methodology well known to those skilled in the art. Specific oligonucleotide probes based upon the nucleic acid sequences of the present invention may be designed and synthesized by methods known in the art. Moreover, the entire sequences may be used directly to synthesize DNA probes by methods known to the skilled artisan such as random primer DNA labelling, nick translation, or end-labelling techniques, or RNA probes using available *in vitro* transcription systems. In addition, specific primers may be designed and used to amplify a part or all of the sequences of the present invention. The resulting amplification products may be labelled directly during amplification reactions or labelled after amplification reactions, and used as probes to isolate full-length cDNA or genomic fragments under conditions of appropriate stringency.

In addition, short segments of the nucleic acids or nucleic acid fragments of the present invention may be used in protocols to amplify longer nucleic acids or

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nucleic acid fragments encoding homologous genes from DNA or RNA. For example, polymerase chain reaction may be performed on a library of cloned nucleic acid fragments wherein the sequence of one primer is derived from the nucleic acid sequences of the present invention, and the sequence of the other primer takes advantage of the presence of the polyadenylic acid tracts to the 3' end of the mRNA precursor encoding plant genes. Alternatively, the second primer sequence may be based upon sequences derived from the cloning vector. For example, those skilled in the art can follow the RACE protocol [Frohman et al. (1988), Proc. Natl. Acad. Sci. USA 85:8998, the entire disclosure of which is incorporated herein by reference] to generate cDNAs by using PCR to amplify copies of the region between a single point in the transcript and the 3' or 5' end. Using commercially available 3' RACE and 5' RACE systems (BRL), specific 3' or 5' cDNA fragments may be isolated [Ohara et al. (1989), Proc. Natl. Acad. Sci. USA 86:5673; Loh et al. (1989), Science 243:217, the entire disclosures of which are incorporated herein by reference]. Products generated by the 3' and 5' RACE procedures may be combined to generate full-length cDNAs.

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In a second aspect of the present invention there is provided a substantially purified or isolated polypeptide from a clover, (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species, selected from the group consisting of CHS and CHS-like, BAN and BAN-like, and LAR and LAR-like proteins; and functionally active fragments and variants thereof.

The clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species may be of any suitable type, including white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*), alfalfa (*Medicago sativa*), Italian or annual ryegrass (*Lolium multiflorum*), perennial ryegrass (*Lolium perenne*), tall fescue (*Festuca arundinacea*), meadow fescue (*Festuca pratensis*) and red fescue (*Festuca rubra*). Preferably the species is a clover or a ryegrass, more preferably white clover (*T. repens*) or perennial ryegrass (*L. perenne*).

In a preferred embodiment of this aspect of the invention, the substantially purified or isolated CHS or CHS-like polypeptide includes an amino acid sequence selected from the group consisting of sequences shown in Figures 3, 7, 11 and 15

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hereto (Sequence ID Nos. 2, 4, 6 and 8, respectively) and functionally active fragments and variants thereof.

In a further preferred embodiment of this aspect of the invention, the substantially purified or isolated BAN or BAN-like polypeptide includes an amino acid sequence shown in Figure 19 hereto (Sequence ID No. 10), and functionally active fragments and variants thereof.

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In a still further preferred embodiment of this aspect of the invention, the substantially purified or isolated LAR or LAR-like polypeptide includes an amino acid sequence selected from the group consisting of sequences shown in Figures 23, 27 and 31 hereto (Sequence ID Nos. 12, 14 and 16, respectively), and functionally active fragments and variants thereof.

In a further embodiment of this aspect of the invention, there is provided a polypeptide produced (e.g. recombinantly) from a nucleic acid or nucleic acid fragment according to the present invention. Techniques for recombinantly producing polypeptides are well known to those skilled in the art.

Availability of the nucleotide sequences of the present invention and deduced amino acid sequences facilitates immunological screening of cDNA expression libraries. Synthetic peptides representing portions of the instant amino acid sequences may be synthesized. These peptides may be used to immunise animals to produce polyclonal or monoclonal antibodies with specificity for peptides and/or proteins including the amino acid sequences. These antibodies may be then used to screen cDNA expression libraries to isolate full-length cDNA clones of interest.

In a still further aspect of the present invention there is provided a construct including one or more nucleic acids or nucleic acid fragments according to the present invention.

In a particularly preferred embodiment the construct may include nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides.

In another preferred embodiment the construct may include nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides.

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In yet another preferred embodiment the construct may include nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides.

In an even more preferred embodiment the construct may include nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides.

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Constructs including nucleic acids or nucleic acid fragments encoding CHS or CHS-like and BAN or BAN-like, and optionally further including nucleic acids or nucleic acid fragments encoding LAR or LAR-like, are particularly preferred.

In a still further aspect of the present invention there is provided a vector including one or more nucleic acids or nucleic acid fragments according to the present invention.

In a preferred embodiment of this aspect of the invention, the construct may include one or several of the following: one or more regulatory elements such as promoters, one or more nucleic acids or nucleic acid fragments according to the present invention and one or more terminators; said one or more regulatory elements, one or more nucleic acids or nucleic acid fragments and one or more terminators being operatively linked.

In a particularly preferred embodiment the construct may contain nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides, operatively linked to a regulatory element or regulatory elements, such that both CHS or CHS-like and BAN or BAN-like polypeptides are expressed.

In another preferred embodiment the construct may contain nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides, operatively linked to a regulatory element or regulatory elements, such that both CHS or CHS-like and LAR or LAR-like polypeptides are expressed.

In yet another preferred embodiment the construct may contain nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides, operatively linked to a regulatory element or regulatory elements, such that both LAR or LAR-like and BAN or BAN-like polypeptides are expressed.

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In an even more preferred embodiment the construct may contain nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides, operatively linked to a regulatory element or regulatory elements, such that all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides are expressed.

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Constructs including nucleic acids or nucleic acid fragments encoding CHS or CHS-like and BAN or BAN-like, and optionally further including nucleic acids or nucleic acid fragments encoding LAR or LAR-like, are particularly preferred.

The construct or vector may be of any suitable type and may be viral or non-viral. The vector may be an expression vector. Such vectors include chromosomal, non-chromosomal and synthetic nucleic acid sequences, e.g. derivatives of plant viruses; bacterial plasmids; derivatives of the Ti plasmid from *Agrobacterium tumefaciens*, derivatives of the Ri plasmid from *Agrobacterium rhizogenes*; phage DNA; yeast artificial chromosomes; bacterial artificial chromosomes; binary bacterial artificial chromosomes; vectors derived from combinations of plasmids and phage DNA. However, any other vector may be used as long as it is replicable, integrative or viable in the plant cell.

The regulatory element and terminator may be of any suitable type and may be endogenous to the target plant cell or may be exogenous, provided that they are functional in the target plant cell.

Preferably the regulatory element is a promoter. A variety of promoters which may be employed in the vectors of the present invention are well known to those skilled in the art. Factors influencing the choice of promoter include the desired tissue specificity of the vector, and whether constitutive or inducible expression is desired and the nature of the plant cell to be transformed (e.g. monocotyledon or dicotyledon). Particularly suitable promoters include but are not limited to the constitutive Cauliflower Mosaic Virus 35S (CaMV 35S) promoter and derivatives thereof, the maize Ubiquitin promoter, the rice Actin promoter, and the tissue-specific Arabidopsis small subunit (ASSU) promoter.

A variety of terminators which may be employed in the vectors and constructs of the present invention are also well known to those skilled in the art. The terminator may be from the same gene as the promoter sequence or a

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different gene. Particularly suitable terminators are polyadenylation signals, such as the CaMV 35S polyA and other terminators from the nopaline synthase (*nos*), the octopine synthase (*ocs*) and the rbcS genes.

The construct or vector, in addition to the regulatory element(s), the nucleic acid(s) or nucleic acid fragment(s) of the present invention and the terminator(s), may include further elements necessary for expression of the nucleic acid(s) or nucleic acid fragment(s), in different combinations, for example vector backbone, origin of replication (ori), multiple cloning sites, recognition sites for recombination events, spacer sequences, enhancers, introns (such as the maize Ubiquitin *Ubi* intron), antibiotic resistance genes and other selectable marker genes [such as the neomycin phosphotransferase (npt2) gene, the hygromycin phosphotransferase (hph) gene, the phosphinotricin acetyltransferase (bar or pat) gene and the gentamycin acetyl transferase (aacC1) gene], and reporter genes [such as beta-glucuronidase (GUS) gene (gusA) and green fluorescent protein (gfp)]. The vector may also contain a ribosome binding site for translation initiation. The vector may also include appropriate sequences for amplifying expression.

As an alternative to use of a selectable marker gene to provide a phenotypic trait for selection of transformed host cells, the presence of the vector in transformed cells may be determined by other techniques well known in the art, such as PCR (polymerase chain reaction), Southern blot hybridisation analysis, histochemical GUS assays, visual examination including microscopic examination of fluorescence emitted by gfp, northern and Western blot hybridisation analyses.

Those skilled in the art will appreciate that the various components of the construct or vector are operatively linked, so as to result in expression of said nucleic acid(s) or nucleic acid fragment(s). Techniques for operatively linking the components of the vector of the present invention are well known to those skilled in the art. Such techniques include the use of linkers, such as synthetic linkers, for example including one or more restriction enzyme sites.

The constructs and vectors of the present invention may be incorporated into a variety of plants, including monocotyledons (such as grasses from the genera *Lolium*, *Festuca*, *Paspalum*, *Pennisetum*, *Panicum* and other forage and turfgrasses, corn, oat, sugarcane, wheat and barley), dicotyledons (such as *Arabidopsis*, tobacco, clovers, medics, eucalyptus, potato, sugarbeet, canola,

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soybean, chickpea) and gymnosperms. In a preferred embodiment, the vectors may be used to transform monocotyledons, preferably grass species such as ryegrasses (*Lolium* species) and fescues (*Festuca* species), more preferably perennial ryegrass, including forage- and turf-type cultivars. In an alternate preferred embodiment, the constructs and vectors may be used to transform dicotyledons, preferably forage legume species such as clovers (*Trifolium* species) and medics (*Medicago* species), more preferably white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*) and alfalfa (*Medicago sativa*). Clovers, alfalfa and medics are key pasture legumes in temperate climates throughout the world.

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Techniques for incorporating the constructs and vectors of the present invention into plant cells (for example by transduction, transfection or transformation) are known to those skilled in the art. Such techniques include *Agrobacterium*-mediated introduction, electroporation to tissues, cells and protoplasts, protoplast fusion, injection into reproductive organs, injection into immature embryos and high velocity projectile introduction to cells, tissues, calli, immature and mature embryos. The choice of technique will depend largely on the type of plant to be transformed.

In a further aspect of the present invention there is provided a method of isogenic transformation of a dicotyledonous plant, said method including transforming only one of each pair of cotyledons. This enables the production of pairs of transgenic plant and corresponding untransformed negative control in an otherwise isogenic genetic background for detailed functional assessment of the impact of the transgene on plant phenotype. In a preferred embodiment of this aspect of the invention, the method may include isogenic transformation of a dicotyledonous plant with a construct or vector according to the present invention.

Cells incorporating the constructs and vectors of the present invention may be selected, as described above, and then cultured in an appropriate medium to regenerate transformed plants, using techniques well known in the art. The culture conditions, such as temperature, pH and the like, will be apparent to the person skilled in the art. The resulting plants may be reproduced, either sexually or asexually, using methods well known in the art, to produce successive generations of transformed plants.

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In a further aspect of the present invention there is provided a plant cell, plant, plant seed or other plant part, including, e.g. transformed with, one or more constructs, vectors, nucleic acids or nucleic acid fragments of the present invention.

The plant cell, plant, plant seed or other plant part may be from any suitable species, including monocotyledons, dicotyledons and gymnosperms. In a preferred embodiment the plant cell, plant, plant seed or other plant part may be from a monocotyledon, preferably a grass species, more preferably a ryegrass (*Lolium* species) or fescue (*Festuca* species), more preferably perennial ryegrass, including both forage- and turf-type cultivars. In an alternate preferred embodiment the plant cell, plant, plant seed or other plant part may be from a dicotyledon, preferably forage legume species such as clovers (*Trifolium* species) and medics (*Medicago* species), more preferably white clover (*Trifolium repens*), red clover (*Trifolium pratense*), subterranean clover (*Trifolium subterraneum*) and alfalfa (*Medicago sativa*).

The present invention also provides a plant, plant seed or other plant part, or a plant extract derived from a plant cell of the present invention.

The present invention also provides a plant, plant seed or other plant part, or a plant extract derived from a plant of the present invention.

In a further aspect of the present invention there is provided a method of modifying condensed tannin biosynthesis; of modifying flavonoid biosynthesis; of modifying protein binding, metal chelation, anti-oxidation, and UV-light absorption; of modifying plant pigment production; of modifying plant defence to biotic stresses such as viruses, microorganisms, insects, fungal pathogens; of modifying forage quality by disrupting protein foam and conferring protection from rumen pasture bloat, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment, construct and/or vector according to the present invention.

In a particularly preferred embodiment the method may include introducing into said plant nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides.

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In another preferred embodiment the method may include introducing into said plant nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides.

In yet another preferred embodiment the method may include introducing into said plant nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides.

In an even more preferred embodiment the method may include introducing into said plant nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides.

Methods including the combinatorial expression of nucleic acids or nucleic acid fragments encoding CHS or CHS-like and BAN or BAN-like, and optionally further including the use of nucleic acids or nucleic acid fragments encoding LAR or LAR-like, are particularly preferred.

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In a further aspect of the present invention there is provided a method of inhibiting bloat in an animal, said method including providing the animal with a forage plant including a construct, vector, nucleic acid or nucleic acid fragment according to the present invention. The animal is preferably a ruminant, including sheep, goats and cattle. The forage plant including a construct vector, nucleic acid or nucleic acid fragment according to the present invention may form all or part of the feed of the animal. The forage plant preferably expresses CHS or CHS-like proteins, BAN or BAN-like proteins, and/or LAR or LAR-like proteins at higher levels than the equivalent wild-type plant. More preferably, the forage plant expresses both CHS or CHS-like proteins and BAN or BAN-like proteins; both CHS or CHS-like proteins and LAR or LAR-like proteins; or both BAN or BAN-like proteins and LAR or LAR-like proteins; at higher levels than the equivalent wild-type plant. More preferably, the forage plant expresses all three of CHS or CHS-like proteins, BAN or BAN-like proteins, and LAR or LAR-like proteins; at higher levels than the equivalent wild-type plant.

In a further aspect of the present invention there is provided a method for enhancing an animal's growth rate, said method including providing the animal with a forage plant including a construct, vector, nucleic acid or nucleic acid fragment according to the present invention. The animal is preferably a ruminant,

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including sheep, goats and cattle. The forage plant including a construct, vector, nucleic acid or nucleic acid fragment according to the present invention may form all or part of the feed of the animal. The forage plant preferably expresses CHS or CHS-like proteins, BAN or BAN-like proteins, and/or LAR or LAR-like proteins at higher levels than the equivalent wild-type plant. More preferably, the forage plant expresses both CHS or CHS-like proteins and BAN or BAN-like proteins; both CHS or CHS-like proteins and LAR or LAR-like proteins; or both BAN or BAN-like proteins and LAR or LAR-like proteins; at higher levels than the equivalent wild-type plant. More preferably, the forage plant expresses all three of CHS or CHS-like proteins, BAN or BAN-like proteins, and LAR or LAR-like proteins; at higher levels than the equivalent wild-type plant.

It is estimated that the method of enhancing an animal's growth rate according to this invention should result in an increase in, for example, lamb growth rate of at least approximately 5%, more preferably at least approximately 10%.

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Using the methods and materials of the present invention, condensed tannin biosynthesis, flavonoid biosynthesis, protein binding, metal chelation, antioxidation, UV-light absorption, tolerance to biotic stresses such as viruses, microorganisms, insects and fungal pathogens; pigmentation in for example flowers and leaves; herbage quality and bloat-safety; isoflavonoid content leading to health benefits, may be increased or otherwise altered, for example by incorporating additional copies of one or more sense nucleic acids or nucleic acid fragments of the present invention. They may be decreased or otherwise altered, for example by incorporating one or more antisense nucleic acids or nucleic acid fragments of the present invention.

Documents cited in this specification are for reference purposes only and their inclusion is not acknowledgment that they form part of the common general knowledge in the relevant art.

The present invention will now be more fully described with reference to the accompanying Examples and drawings. It should be understood, however, that the description following is illustrative only and should not be taken in any way as a restriction on the generality of the invention described above.

In the Figures

Figure 1 shows the plasmid map in pGEM-T Easy of TrCHSa3.

Figure 2 shows the nucleotide sequence of TrCHSa3 (Sequence ID No. 1).

Figure 3 shows the deduced amino acid sequence of TrCHSa3 (Sequence ID No.

5 2).

Figure 4 shows plasmid maps of sense and antisense constructs of TrCHSa3 in the binary vector pPZP221:35S<sup>2</sup>.

Figure 5 shows the plasmid map in pGEM-T Easy of TrCHSc.

Figure 6 shows the nucleotide sequence of TrCHSc (Sequence ID No. 3).

10 Figure 7 shows the deduced amino acid sequence of TrCHSc (Sequence ID No. 4).

Figure 8 shows plasmid maps of sense and antisense constructs of TrCHSc in the binary vector pPZP221:35S<sup>2</sup>.

Figure 9 shows the plasmid map in pGEM-T Easy of TrCHSf.

15 Figure 10 shows the nucleotide sequence of TrCHSf (Sequence ID No. 5).

Figure 11 shows the deduced amino acid sequence of TrCHSf (Sequence ID No. 6).

Figure 12 shows plasmid maps of sense and antisense constructs of TrCHSf in the binary vector pPZP221:35S<sup>2</sup>.

20 Figure 13 shows the plasmid map in pGEM-T Easy of TrCHSh.

Figure 14 shows the nucleotide sequence of TrCHSh (Sequence ID No. 7).

Figure 15 shows the deduced amino acid sequence of TrCHSh (Sequence ID No. 8).

Figure 16 shows plasmid maps of sense and antisense constructs of TrCHSh in the binary vector pPZP221:35S<sup>2</sup>.

Figure 17 shows the plasmid map in pGEM-T Easy of TrBANa.

Figure 18 shows the nucleotide sequence of TrBANa (Sequence ID No. 9).

Figure 19 shows the deduced amino acid sequence of TrBANa (Sequence ID No. 10).

Figure 20 shows plasmid maps of sense and antisense constructs TrBANa in the binary vector pPZP221:35S<sup>2</sup>.

5 Figure 21 shows the plasmid map in pGEM-T Easy of TrLARa.

Figure 22 shows the nucleotide sequence of TrLARa (Sequence ID No. 11).

Figure 23 shows the deduced amino acid sequence of TrLARa (Sequence ID No. 12).

Figure 24 shows plasmid maps of sense and antisense constructs of TrLARa in the binary vector pPZP221:35S<sup>2</sup>.

Figure 25 shows the plasmid map in pGEM-T Easy of TrLARb.

Figure 26 shows the nucleotide sequence of TrLARb (Sequence ID No. 13).

Figure 27 shows the deduced amino acid sequence of TrLARb (Sequence ID No. 14).

Figure 28 shows plasmid maps of sense and antisense constructs of TrLARb in the binary vector pPZP221:35S<sup>2</sup>.

Figure 29 shows the plasmid map in pGEM-T Easy of TrLARc.

Figure 30 shows the nucleotide sequence of TrLARc (Sequence ID No. 15).

Figure 31 shows the deduced amino acid sequence of TrLARc (Sequence ID No. 20 16).

Figure 32 shows plasmid maps of sense and antisense constructs of TrLARc in the binary vector pPZP221:35S<sup>2</sup>.

Figure 33 shows the plasmid map of the binary vector pPZP221:ASSU::TrBAN:35S<sup>2</sup>::TrCHS.

25 Figure 34 shows the plasmid maps of the modular vector system comprising a binary base vector and 7 auxiliary vectors.

Figure 35 shows an example of the modular binary transformation vector system comprising plasmid maps of the binary transformation vector backbone and 4

expression cassettes in auxiliary vectors (A) and the plasmid map of the T-DNA region of the final binary transformation vector.

Figure 36 shows A, white clover cotyledons; B, C, D, selection of plantlets transformed with a binary transformation vector constructed as described in Examples 4 and 5; E, putative transgenic white clover on root-inducing medium; F, G, white clover plants transgenic for genes involved in condensed tannin biosynthesis.

Figure 37 shows the molecular analysis of white clover plants transgenic for the TrBAN gene with Q-PCR amplification plot, agarose gel of PCR product and Southern hybridisation blot.

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Figure 38 shows the molecular analysis of white clover plants transgenic for the TrCHSf gene with Q-PCR amplification plot and agarose gel of PCR product.

Figure 39 shows the molecular analysis of white clover plants transgenic for the TrLARb gene with Q-PCR amplification plot, agarose gel of PCR product and Southern hybridisation blot.

#### **EXAMPLE 1**

# Preparation of cDNA libraries, isolation and sequencing of cDNAs coding for CHS, CHS-like, BAN, BAN-like, LAR and LAR-like proteins from white clover (*Trifolium repens*)

cDNA libraries representing mRNAs from various organs and tissues of white clover (*Trifolium repens*) were prepared. The characteristics of the white clover libraries are described below (Table 1).

TABLE 1 cDNA libraries from white clover (*Trifolium repens*)

Library	Organ/Tissue						
01wc	Whole seedling, light grown						
02wc	Nodulated root 3, 5, 10, 14, 21 &28 day old seedling						
03wc	Nodules pinched off roots of 42 day old rhizobium inoculated plants						
04wc	Cut leaf and stem collected after 0, 1, 4, 6 &14 h after cutting						
05wc	Inflorescences: <50% open, not fully open and fully open						

Library	Organ/Tissue
06wc	Dark grown etiolated
07wc	Inflorescence – very early stages, stem elongation, < 15 petals, 15-20 petals
08wc	seed frozen at -80°C, imbibed in dark overnight at 10°C
09wc	Drought stressed plants
10wc	AMV infected leaf
11wc	WCMV infected leaf
12wc	Phosphorus starved plants
13wc	Vegetative stolon tip
14wc	stolon root initials
15wc	Senescing stolon
16wc	Senescing leaf

The cDNA libraries may be prepared by any of many methods available. For example, total RNA may be isolated using the Trizol method (Gibco-BRL, USA) or the RNeasy Plant Mini kit (Qiagen, Germany), following the manufacturers' instructions. cDNAs may be generated using the SMART PCR cDNA synthesis kit (Clontech, USA), cDNAs may be amplified by long distance polymerase chain reaction using the Advantage 2 PCR Enzyme system (Clontech, USA), cDNAs may be cleaned using the GeneClean spin column (Bio 101, USA), tailed and size fractionated, according to the protocol provided by Clontech. The cDNAs may be introduced into the pGEM-T Easy Vector system 1 (Promega, USA) according to the protocol provided by Promega. The cDNAs in the pGEM-T Easy plasmid vector are transfected into *Escherichia coli* Epicurean coli XL10-Gold ultra competent cells (Stratagene, USA) according to the protocol provided by Stratagene.

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Alternatively, the cDNAs may be introduced into plasmid vectors for first preparing the cDNA libraries in Uni-ZAP XR vectors according to the

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manufacturer's protocol (Stratagene Cloning Systems, La Jolla, CA, USA). The Uni-ZAP XR libraries are converted into plasmid libraries according to the protocol provided by Stratagene. Upon conversion, cDNA inserts will be contained in the plasmid vector pBlueScript. In addition, the cDNAs may be introduced directly into precut pBlueScript II SK(+) vectors (Stratagene) using T4 DNA ligase (New England Biolabs), followed by transfection into *E. coli* DH10B cells according to the manufacturer's protocol (GIBCO BRL Products).

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Once the cDNA inserts are in plasmid vectors, plasmid DNAs are prepared from randomly picked bacterial colonies containing recombinant plasmids, or the insert cDNA sequences are amplified via polymerase chain reaction using primers specific for vector sequences flanking the inserted cDNA sequences. Plasmid DNA preparation may be performed robotically using the Qiagen QiaPrep Turbo kit (Qiagen, Germany) according to the protocol provided by Qiagen. Amplified insert DNAs are sequenced in dye-terminator sequencing reactions to generate partial cDNA sequences (expressed sequence tags or "ESTs"). The resulting ESTs are analysed using an Applied Biosystems ABI 3700 sequence analyser.

## EXAMPLE 2 DNA sequence analyses

The cDNA clones encoding CHS, CHS-like, BAN, BAN-like, LAR and LAR-like proteins were identified by conducting BLAST (Basic Local Alignment Search Tool; Altschul *et al.* (1993), *J. Mol. Biol.* 215:403-410) searches. The cDNA sequences obtained were analysed for similarity to all publicly available DNA sequences contained in the eBioinformatics nucleotide database using the BLASTN algorithm provided by the National Center for Biotechnology Information (NCBI). The DNA sequences were translated in all reading frames and compared for similarity to all publicly available protein sequences contained in the SWISS-PROT protein sequence database using BLASTx algorithm (v 2.0.1) (Gish and States (1993), *Nature Genetics* 3:266-272) provided by the NCBI.

The cDNA sequences obtained and identified were then used to identify additional identical and/or overlapping cDNA sequences generated using the BLASTN algorithm. The identical and/or overlapping sequences were subjected to a multiple alignment using the CLUSTALw algorithm, and to generate a consensus contig sequence derived from this multiple sequence alignment. The

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consensus contig sequence was then used as a query for a search against the SWISS-PROT protein sequence database using the BLASTx algorithm to confirm the initial identification.

#### **EXAMPLE 3**

# 5 Identification and full-length sequencing of cDNAs encoding white clover CHS, BAN and LAR proteins

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To fully characterise for the purposes of the generation of probes for hybridisation experiments and the generation of transformation vectors, a set of cDNAs encoding white clover CHS, BAN and LAR proteins was identified and fully sequenced.

Full-length cDNAs were identified from our EST sequence database using relevant published sequences (NCBI databank) as queries for BLAST searches. Full-length cDNAs were identified by alignment of the query and hit sequences using Sequencher (Gene Codes Corp., Ann Arbor, MI 48108, USA). The original plasmid was then used to transform chemically competent XL-1 cells (prepared inhouse, CaCl<sub>2</sub> protocol). After colony PCR (using HotStarTaq, Qiagen) a minimum of three PCR-positive colonies per transformation were picked for initial sequencing with M13F and M13R primers. The resulting sequences were aligned with the original EST sequence using Sequencher to confirm identity and one of the three clones was picked for full-length sequencing, usually the one with the best initial sequencing result.

Sequencing of TrBAN could be completed with M13F and M13R primers. Sequencing of TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrLARa, TrLARb and TrLARc was completed by primer walking, i.e. oligonucleotide primers were designed to the initial sequence and used for further sequencing. The sequences of the oligonucleotide primers are shown in Table 2.

Contigs were then assembled in Sequencher. The contigs include the sequences of the SMART primers used to generate the initial cDNA library as well as pGEM-T Easy vector sequence up to the EcoRI cut site both at the 5' and 3' end.

Plasmid maps and the full cDNA sequences of TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc proteins were obtained (Figures 1, 2, 5, 6, 9, 10, 13, 14, 17, 18, 21, 22, 25, 26, 29 and 30).

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TABLE 2
List of primers used for sequencing of the full-length cDNAs of TrCHSa3,
TrCHSc, TrCHSf, TrCHSh, TrLARa, TrLARb and TrLARc

gene name	clone ID	sequencing primer	primer sequence (5'>3')
TrCHSa3	05wc1RsB06	05wc1RsB06.f1	AGGAGGCTGCAGTCAAGG
		05wc1RsB06.f2	TGCCTGAAATTGAGAAACC
		05wc1RsB06.f3	AAAGCTAGCCTTGAAGCC
TrCHSc	07wc1TsE12	07wc1TsE12.f1	TCGGACATAACTCATGTGG
		07wc1TsE12.f2	TTGGGTTGGAGAATAAGG
		07wc1TsE12.r1	TGGACATTTATTGGTTGC
		07wc1TsE12.r2	TATCATGTCTGGAAATGC
TrCHSf	07wc1UsD07	07wc1UsD07.f1	AGATTGCATCAAAGAATGG
		07wc1UsD07.r1	GGTCCAAAAGCCAATCC
TrCHSh	13wc2lsG04	13wc2lsG04.f1	TAAGACGAGACATAGTGG
		13wc2lsG04.r1	TATTCACTAAGCACATGC
TrLARa	05wc1CsA02	05wc1CsA02.f1	TCATTTCTGCAATAGGAGG
		05wc1CsA02.r1	ATCCACCTCAGGTGAACC
TrLARb	05wc3EsA03	05wc3EsA03.f1	AATAGGAGGCTCTGATGG
		05wc3EsA03r1	ATCCACCTCAGGTGAACC
TrLARc	07wc1VsF06	07wc1VsF06.f1	AGGCTCTGATGGCTTGC
		07wc1VsF06.r1	ATCCACCTCAGGTGAACC

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#### **EXAMPLE 4**

Development of binary transformation vectors containing chimeric genes with cDNA sequences from white clover TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc

To alter the expression of the proteins involved in flavonoid biosynthesis, and more specifically condensed tannin biosynthesis to improve herbage quality

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and bloat-safety, a set of sense and antisense binary transformation vectors was produced.

cDNA fragments were generated by high fidelity PCR with a proofreading DNA polymerase using the original pGEM-T Easy plasmid cDNA as a template. The primers used (Table 3) contained recognition sites for appropriate restriction enzymes, for example EcoRI and Xbal, for directional and non-directional cloning into the target vector. After PCR amplification and restriction digest with the appropriate restriction enzyme (usually Xbal), the cDNA fragments were cloned into the corresponding site in a modified pPZP binary vector (Hajdukiewicz et al., 1994). The pPZP221 vector was modified to contain the 35S2 cassette from pKYLX71:35S<sup>2</sup> (Schardl et al., 1987) as follows: pKYLX71:35S<sup>2</sup> was cut with Clal. The 5' overhang was filled in using Klenow and the blunt end was A-tailed with Tag polymerase. After cutting with EcoRI, the 2kb fragment with an EcoRIcompatible and a 3'-A tail was gel-purified. pPZP221 was cut with HindIII and the resulting 5' overhang filled in and T-tailed with Tag polymerase. The remainder of the original pPZP221 multi-cloning site was removed by digestion with EcoRI, and the expression cassette cloned into the EcoRI site and the 3' T overhang restoring the HindIII site. This binary vector contains between the left and right border the plant selectable marker gene aacC1 under the control of the 35S promoter and 35S terminator and the pKYLX71:35S2-derived expression cassette with a CaMV 35S promoter with a duplicated enhancer region and an rbcS terminator.

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Alternatively, the primers for the amplification of cDNA fragments contained attB sequences for use with recombinases utilising the GATEWAY® system (Invitrogen). The resulting PCR fragments were used in a recombination reaction with pDONR® vector (Invitrogen) to generate entry vectors. A GATEWAY® cloning cassette (Invitrogen) was introduced into the multicloning site of the pPZP221:35S² vector following the manufacturer's protocol. In a further recombination reaction, the cDNAs encoding the open reading frame sequences were transferred from the entry vector to the GATEWAY®-enabled pPZP221:35S² vector.

The orientation of the constructs (sense or antisense) was checked by restriction enzyme digest and sequencing which also confirmed the correctness of the sequence. Transformation vectors containing chimeric genes using full-length

open reading frame cDNAs encoding white clover TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc proteins in sense and antisense orientation under the control of the CaMV 35S<sup>2</sup> promoter were generated (Figures 4, 8, 12, 16, 20, 24, 28 and 32).

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TABLE 3

List of primers used to PCR-amplify the open reading frames

gene name	primer	primer sequence (5'->3')
TrCHSa3	05wc1RsB06f	GAATTCTAGAAGATATGGTGAGTGTAGCTG
	05wc1RsB06r	GAATTCTAGAATCACACATCTTATATAGCC
TrCHSa3	05wc1RsB06fG	GGGGACAAGTTTGTACAAAAAAGCAGGCTTCTAGA AGATATGGTGAGTGTAGCTG
	05wc1RsB06rG	GGGGACCACTTTGTACAAGAAAGCTGGGTTCTAGA ATCACACATCTTATATAGCC
TrCHSc	07wc1TsE12f	GAATTCTAGAAGAAGAAATATGGGAGACGAAGG
	07wc1TsE12r	GAATTCTAGAAAGACTTCATGCACACAAGTTCC
TrCHSf	07wc1UsD07f	GAATTCTAGATGATTCATTGTTTGTTTCCATAAC
	07wc1UsD07r	GAATTCTAGAACATATTCATCTTCCTATCAC
TrCHSh	13wc2lsG04f	GAATTCTAGATCCAAATTCTCGTACCTCACC
	13wc2lsG04r	GAATTCTAGATAGTTCACATCTCTCGGCAGG
TrBANa	05wc2XsG02f	GGATCCTCTAGAGCACTAGTGTGTATAAGTTTCTT GG
	05wc2XsG02r	GGATCCTCTAGACCCCCTTAGTCTTAAAATACTCG
TrLARa	05wc1CsA02fG	GGGGACAAGTTTGTACAAAAAAGCAGGCTCTAGAA AGCAAAGCAA
	05wc1CsA02rG	GGGGACCACTTTGTACAAGAAAGCTGGGTCTAGAT CCACCTCAGGTGAACC
TrLARb	05wc3EsA03fG	GGGGACAAGTTTGTACAAAAAAGCAGGCTCTAGAA AGCAATGGCACCAGCAGC
	05wc3EsA03rG	GGGGACCACTTTGTACAAGAAAGCTGGGTCTAGAT CCACCTCAGGTGAACC
TrLARc	07wc1VsF06fG	GGGGACAAGTTTGTACAAAAAAGCAGGCTCTAGAT AAAGCAATGGCACCAGC
	07wc1VsF06rG	GGGGACCACTTTGTACAAGAAAGCTGGGTCTAGAT CCACCTCAGGTGAACC

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The pPZP221:35S<sup>2</sup> binary vector was further modified to contain two expression cassettes within one T-DNA. The expression cassette from the binary vector pWM5 consisting of the ASSU promoter and the tob terminator was PCR-amplified with a proofreading DNA polymerase using oligonucleotide primers with the following sequences:

forward primer 5'-CCACCATGTTTGAAATTTATTATGTGTTTTTTCCG-3';
reverse primer 5'-TAATCCCGGGTAAGGGCAGCCCATACAAATGAAGC-3'.

The PCR product was cut with BstXI and Smal and cloned directionally into the equally cut pPZP221:35S² vector. Additionally, a GATEWAY® cloning cassette (Invitrogen) was introduced into the multicloning site in the 35S²:rbcS expression cassette following the manufacturer's protocol. TrBANa was cloned into the ASSU:tob expression cassette, TrCHSa3 was amplified with the GATEWAY®-compatible primers (see Table 3) and cloned into the 35S²:rbcS expression cassette. A transformation vector containing chimeric genes using full-length open reading frame cDNAs encoding white clover TrBANa protein in sense orientation under the control of the ASSU promoter and TrCHSc3 protein in sense orientation under the control of the CaMV 35S² promoter within the same T-DNA was generated (Figure 33).

20 EXAMPLE 5

Development of binary transformation vectors containing chimeric genes with a combination of 2 or more cDNA sequences from white clover TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc

To alter the expression of the proteins involved in flavonoid biosynthesis, and more specifically condensed tannin biosynthesis to improve herbage quality and bloat-safety, a modular binary transformation vector system was used (Figure 34). The modular binary vector system enables simultaneous integration of up to seven transgenes the expression of which is controlled by individual promoter and terminator sequences into the plant genome (Goderis *et al.*, 2002).

The modular binary vector system consists of a pPZP200-derived vector (Hajdukiewicz et al., 1994) backbone containing within the T-DNA a number of unique restriction sites recognised by homing endonucleases. The same

restriction sites are present in pUC18-based auxiliary vectors flanking standard multicloning sites. Expression cassettes comprising a selectable marker gene sequence or a cDNA sequence to be introduced into the plant under the control of regulatory sequences like promoter and terminator can be constructed in the auxiliary vectors and then transferred to the binary vector backbone utilising the homing endonuclease restriction sites. Up to seven expression cassettes can thus be integrated into a single binary transformation vector. The system is highly flexible and allows for any combination of cDNA sequence to be introduced into the plant with any regulatory sequence.

For example, a selectable marker cassette comprising the nos promoter and nos terminator regulatory sequences controlling the expression of the nptll gene was PCR-amplified using a proofreading DNA polymerase from the binary vector pKYLX71:35S<sup>2</sup> and directionally cloned into the Agel and Notl sites of the auxiliary vector pAUX3166. Equally, other selectable marker cassettes can be introduced into any of the auxiliary vectors. 15

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In another example, the expression cassette from the binary vector pWM5 consisting of the ASSU promoter and the tob terminator was PCR-amplified with a proofreading DNA polymerase and directionally cloned into the Agel and Notl sites of the auxiliary vector pAUX3169. Equally, other expression cassettes can be introduced into any of the auxiliary vectors.

In yet another example, the expression cassette from the direct gene transfer vector pDH51 was cut using EcoRI and cloned directly into the EcoRI site of the auxiliary vector pAUX3132.

TABLE 4

List of primers used to PCR-amplify plant selectable marker cassettes and 25 the regulatory elements used to control the expression of TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc genes

expression cassette	primer	primer sequence (5'>3')
nos::nptII-nos	forward	ATAATAACCGGTTGATCATGAGCGGAGAATTAAG GG
	reverse	ATAATAGCGGCCGCTAGTAACATAGATGACACCG CG

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expression cassette	primer	primer sequence (5'>3')
35S::aacC1-35S	forward	AATAGCGGCCGCGATTTAGTACTGGATTTTGG
	reverse	AATAACCGGTACCCACGAAGGAGCATCGTGG
35S <sup>2</sup> ::rbcS	forward	ATAATAACCGGTGCCCGGGGATCTCCTTTGCC
	reverse	ATAATAGCGGCCGCATGCATGTTGTCAATCAATT GG
assu::tob	forward	TAATACCGGTAAATTTATTATGRGTTTTTTTCCG
	reverse	TAATGCGGCCGCTAAGGGCAGCCCATACAAATGA AGC

The expression cassettes were further modified by introducing a GATEWAY® cloning cassette (Invitrogen) into the multicloning site of the respective pAUX vector following the manufacturer's protocol. In a recombination reaction, the cDNAs encoding the open reading frame sequences were transferred from the entry vector obtained as described in Example 4 to the GATEWAY®-enabled pAUX vector. Any combination of the regulatory elements with cDNA sequences of TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc can be obtained. One typical example is given in Figure 35 with expression cassettes for the nptII plant selectable marker, TrBANa, TrLARa and TrCHSa3.

Complete expression cassettes comprising any combination of regulatory elements and cDNA sequences to be introduced into the plant were then cut from the auxiliary vectors using the respective homing endonuclease and cloned into the respective restriction site on the binary vector backbone. After verification of the construct by nucleotide sequencing, the binary transformation vector comprising a number of expression cassettes was used to generate transgenic white clover plants.

#### **EXAMPLE 6**

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Production by Agrobacterium-mediated transformation and analysis of transgenic white clover plants carrying chimeric white clover TrCHSa3, TrCHSc, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc genes involved in flavonoid biosynthesis

A set of binary transformation vectors carrying chimeric white clover genes involved in flavonoid biosynthesis, and more specifically condensed tannin biosynthesis to improve herbage quality and bloat-safety, were produced as detailed in Examples 4 and 5.

Agrobacterium-mediated gene transfer experiments were performed using these transformation vectors.

The production of transgenic white clover plants carrying the white clover TrCHSa3, TrCHSc, TrCHSf, TrCHSh, TrBANa, TrLARa, TrLARb and TrLARc cDNAs, either singly or in combination, is described here in detail.

#### Preparation of Agrobacterium

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Agrobacterium tumefaciens strain AGL-1 transformed with one of the binary vector constructs detailed in Example 6 were streaked on LB medium containing 50 μg/ml rifampicin and 50 μg/ml kanamycin and grown at 27 °C for 48 hours. A single colony was used to inoculate 5 ml of LB medium containing 50 μg/ml rifampicin and 50 μg/ml kanamycin and grown over night at 27 °C and 250 rpm on an orbital shaker. The overnight culture was used as an inoculum for 500 ml of LB medium containing 50 μg/ml kanamycin only. Incubation was over night at 27 °C and 250 rpm on an orbital shaker in a 2 l Erlenmeyer flask.

#### Preparation of white clover seeds

1 spoon of seeds (ca. 500) was placed into a 280 μm mesh size sieve and washed for 5 min under running tap water, taking care not to wash seeds out of sieve. In a laminar flow hood, seeds were transferred with the spoon into an autoclaved 100 ml plastic culture vessel. A magnetic stirrer (wiped with 70% EtOH) and ca. 30 ml 70% EtOH were added, and the seeds were stirred for 5 min. The EtOH was discarded and replaced by 50 ml 1.5% sodium hypochlorite. The seeds were stirred for an additional 45 - 60 min on a magnetic stirrer. The sodium hypochlorite was then discarded and the seeds rinsed 3 to 4 times with autoclaved

ddH<sub>2</sub>O. Finally 30 ml of ddH<sub>2</sub>O were added, and seeds incubated over night at 10 - 15°C in an incubator.

### Agrobacterium-mediated transformation of white clover

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The seed coat and endosperm layer of the white clover seeds prepared as above were removed with a pair of 18 G or 21 G needles. The cotyledons were cut from the hypocotyl leaving a ca. 1.5 mm piece of the cotyledon stalk. The cotyledons were transferred to a petridish containing ddH<sub>2</sub>O. After finishing the isolation of clover cotyledons, ddH<sub>2</sub>O in the petridish was replaced with Agrobacterium suspension (diluted to an  $OD_{600} = 0.2 - 0.4$ ). The petridish was sealed with its lid and incubated for 40 min at room temperature.

After the incubation period, each cotyledon was transferred to paper towel using the small dissection needle, dried and plated onto regeneration medium RM73. The plates were incubated at  $25^{\circ}$ C with a 16h light/8h dark photoperiod. On day 4, the explants were transferred to fresh regeneration medium. Cotyledons transformed with *Agrobacterium* were transferred to RM73 containing cefotaxime (antibacterial agent) and gentamycin. The dishes were sealed with Parafilm and incubated at  $25^{\circ}$ C under a 16/8 h photoperiod. Explants were subcultured every three weeks for a total of nine weeks onto fresh RM 73 containing cefotaxime and gentamycin. Shoots with a green base were then transferred to root-inducing medium RIM. Roots developed after 1-3 weeks, and plantlets were transferred to soil when the roots were well established.

This process is shown in detail in Figure 36.

# Preparation of genomic DNA for real-time PCR and analysis for the presence of transgenes

3 – 4 leaves of white clover plants regenerated on selective medium were harvested and freeze-dried. The tissue was homogenised on a Retsch MM300 mixer mill, then centrifuged for 10 min at 1700xg to collect cell debris. Genomic DNA was isolated from the supernatant using Wizard Magnetic 96 DNA Plant System kits (Promega) on a Biomek FX (Beckman Coulter). 5 μl of the sample (50 μl) were then analysed on an agarose gel to check the yield and the quality of the genomic DNA.

Genomic DNA was analysed for the presence of the transgene by real-time PCR using SYBR Green chemistry. PCR primer pairs (Table 4) were designed using MacVector (Accelrys) or PrimerExpress (ABI). The forward primer was located within the 35S<sup>2</sup> promoter region and the reverse primer within the transgene to amplify products of approximately 150 - 250 bp as recommended. The positioning of the forward primer within the 35S<sup>2</sup> promoter region guaranteed that endogenous genes in white clover were not detected.

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TABLE 5

List of primers used for Real-time PCR analysis of white clover plants transformed with chimeric white clover genes involved in condensed tannin biosynthesis

construct	primer 1 (forward), 5'->3'	primer 2 (reverse), 5'->3'
pPZP221TrCHSa3	CATTTCATTTGGAGAGGACACGC	AACACGGTTTGGTGGATTTGC
pPZP221TrCHSc	TTGGAGAGGACACGCTGAAATC	ACAAGTTGGTGAGGGAATGCC
pPZP221TrCHSf	CATTTCATTTGGAGAGGACACGC	TCGTTGCCTTTCCCTGAGTAGG
pPZP221TrCHSh	TCATTTGGAGAGGACACGCTG	CGGTCACCATTTTTTTTTTGTTGGAGG
pPZP221TrBANa	TTGGAGAGGACACGCTGAAATC	CAACAAAACCAGTGCCACC
pPZP221TrLARa	ATGACGCACAATCCCACTATCC	AGCCTTAGAAGAGAGAAGAGGTCC
pPZP221TrLARb	ATGACGCACAATCCCACTATCC	AGCCTTAGAAGAGAGAAGAGGTCC
pPZP221TrLARc	ATGACGCACAATCCCACTATCC	AGCCTTAGAAGAGAGAAGAGGTCC

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5 μl of each genomic DNA sample was run in a 50 μl PCR reaction including SYBR Green on an ABI 7700 (Applied Biosystems) together with samples containing DNA isolated from wild type white clover plants (negative control), samples containing buffer instead of DNA (buffer control) and samples containing the plasmid used for transformation (positive plasmid control). Cycling conditions used were 2 min. at 50 °C, 10 min. at 95 °C and then 40 cycles of 15 sec. at 95 °C, 1 min. at 60 °C.

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### Preparation of genomic DNA and analysis of DNA for presence and copy number of transgene by Southern hybridisation blotting

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Genomic DNA for Southern hybridisation blotting was obtained from leaf material of white clover plants following the CTAB method. Southern hybridisation blotting experiments were performed following standard protocols as described in Sambrook *et al.* (1989). In brief, genomic DNA samples were digested with appropriate restriction enzymes and the resulting fragments separated on an agarose gel. After transfer to a membrane, a cDNA fragment representing a transgene or selectable marker gene was used to probe the size-fractionated DNA fragments. Hybridisation was performed with either radioactively labelled probes or using the non-radioactive DIG labelling and hybridisation protocol (Boehringer) following the manufacturer's instructions.

Plants were obtained after transformation with all chimeric constructs and selection on medium containing gentamycin. Details of plant analysis are given in Table 5 and Figures 37, 38 and 39.

TABLE 5

Transformation of white clover with binary transformation vectors comprising cDNAs of white clover genes involved in condensed tannin biosyntheses, selection and molecular analysis of regenerated plants.

Southern copy number	range	n/d	p/u	n/d	p/u	7 1 to 4	p/u	5 1 to 3	p/u
QPCR-positive Sout		23 n	27 n	27 n	30	38	38	47	29 r
Soil		32	41	44	37	20	45	47	32
selection into RIM		135	68	113	6.2	144	88	133	96
cotyledons	transformed	2358	3460	3931	3743	2315	2487	3591	2835
construct	·	pPZP221-35S2::TrCHSa3	pPZP221-35S2::TrCHSc	pPZP221-35S2::TrCHSf	pPZP221-35S2::TrCHSh	pPZP221-35S2::TrBANa	pPZP221-35S2::TrLARa	pPZP221-35S2::TrLARb	pPZP221-35S2::TrLARc

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#### **REFERENCES**

- Altschul, S.F., Gish, W., Miller, W., Myers, E.W., Lipman, D.J. (1990) "Basic local alignment search tool." J. Mol. Biol. **215**, 403-410.
- Frohman, M.A., Dush, M.K., Martin, G.R. (1988) Rapid production of full-length cDNAs from rare transcripts: amplification using a single gene-specific oligonucleotide primer. *Proc. Natl. Acad. Sci. USA* **85**, 8998.
  - Gish, W., States, D.J. (1993) Identification of protein coding regions by database similarity search. Nature Genetics **3**, 266-272.
- Goderis, I., De Bolle, M.F.C., Francois, I., Wouters, P.F.J., Broekaert, W.F., and
  Cammue, B.P.A. (2002) A set of modular plant transformation vectors allowing flexible insertion of up to six expression units. Plant Molecular Biology **50**, 17-27.
  - Hajdukiewicz P, Svab Z, Maliga P. (1994) The small, versatile pPZP family of Agrobacterium binary vectors for plant transformation. Plant Mol Biol. **25**, 989-94.
    - Loh, E.Y., Elliott, J.F., Cwirla, S., Lanier, L.L., Davis, M.M. (1989). Polymerase chain reaction with single-sided specificity: Analysis of T-cell receptor delta chain. *Science* **243**, 217-220.
- Ohara, O., Dorit, R.L., Gilbert, W. (1989). One-sided polymerase chain reaction:

  The amplification of cDNA. *Proc. Natl. Acad Sci USA* **86**, 5673-5677
  - Sambrook, J., Fritsch, E.F., Maniatis, T. (1989). Molecular Cloning. A Laboratory Manual. Cold Spring Harbour Laboratory Press
- Schardl, C.L., Byrd, A.D., Benzion, G., Altschuler, M.A., Hildebrand, D.F., Hunt, A.G. (1987) Design and construction of a versatile system for the expression of foreign genes in plants. Gene **61**, 1-11

Finally, it is to be understood that various alterations, modifications and/or additions may be made without departing from the spirit of the present invention as outlined herein.

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PCT/AU2004/000494

#### **CLAIMS**

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- 1. A substantially purified or isolated nucleic acid or nucleic acid fragment encoding a condensed tannin biosynthetic enzyme selected from the group consisting of chalcone synthase (CHS), CHS-like, dihydroflavonol 4-reductase (BAN), BAN-like, leucoanthocyanidine reductase (LAR) and LAR-like, or a functionally active fragment or variant of such a polypeptide, from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species.
- 2. A nucleic acid or nucleic acid fragment according to Claim 1, wherein said nucleic acid or nucleic acid fragment is from white clover (*Trifolium repens*) or perennial ryegrass (*Lolium perenne*).
- 3. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a CHS polypeptide or CHS-like polypeptide and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 2, 6, 10 and 14 hereto (Sequence ID Nos. 1, 3, 5 and 7, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c) and (d).
- 4. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a BAN polypeptide or BAN-like polypeptide and including a nucleotide sequence selected from the group consisting of (a) sequence shown in Figure 18 hereto (Sequence ID No. 9); (b) complements of the sequence recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c) and (d).
  - 5. A nucleic acid or nucleic acid fragment according to Claim 1, encoding a LAR polypeptide or LAR-like polypeptide and including a nucleotide sequence selected from the group consisting of (a) sequences shown in Figures 22, 26 and 30 hereto (Sequence ID Nos. 11, 13 and 15, respectively); (b) complements of the sequences recited in (a); (c) sequences antisense to the sequences recited in (a) and (b); and (d) functionally active fragments and variants

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of the sequences recited in (a), (b) and (c); and (e) RNA sequences corresponding to the sequences recited in (a), (b), (c) and (d).

- 6. A construct including one or more nucleic acid or nucleic acid fragments according to any one of claims 1 to 5.
- 7. A construct according to claim 6 including nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides.
  - 8. A construct according to claim 6 including nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides.
  - 9. A construct according to claim 6 including nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides.

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- 10. A construct according to claim 6 including nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, LAR or LAR-like and BAN or BAN-like polypeptides.
- 15 11. A construct according to any one of claims 6 to 10 wherein the one or more nucleic acids or nucleic acid fragments are operably linked to one or more regulatory elements, such that the one or more nucleic acids or nucleic acid fragments are expressed.
- 12. A construct according to Claim 11, wherein the one or more regulatory elements include a promoter and a terminator, said promoter, nucleic acid or nucleic acid fragment and terminator being operatively linked.
  - 13. A plant cell, plant, plant seed or other plant part, including a construct according to any one of claims 6 to 12.
- 14. A plant, plant seed or other plant part derived from a plant cell or 25 plant according to Claim 13.
  - 15. A method of modifying one or more of condensed tannin biosynthesis; protein binding, metal chelation; anti oxidation; UV-light absorption; pigment production; or plant defence to a biotic stress; in a plant, said method including introducing into said plant an effective amount of a nucleic acid or nucleic acid fragment according to any one or claims 1 to 5 or a construct according or any one of claims 6 to 12.

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- 16. A method according to claim 15 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides.
- 17. A method according to claim 15 wherein said method includes
   5 introducing into said plant effective amounts of nucleic acids or nucleic acid
   fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides
  - 18. A method according to claim 15 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides.
- 19. A method according to claim 15 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides.
- 20. A method according to any one of claims 15 to 19 wherein the method is modifying plant defence to biotic stress and the biotic stress is selected from the group consisting of viruses, micro-organisms, insects and fungal pathogens.
  - 21. A method of modifying forage quality of a plant by disrupting protein foam and/or conferring protection from rumen pasture bloat, said method including introducing into said plant an effective amount of a nucleic acid fragment according to any one of claims 1 to 5 or a construct according to any one of claims 6 to 12.

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- 22. A method according to claim 21 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and BAN or BAN-like polypeptides.
- 23. A method according to claim 21 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding both CHS or CHS-like and LAR or LAR-like polypeptides
- 24. A method according to claim 21 wherein said method includes 30 introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding both LAR or LAR-like and BAN or BAN-like polypeptides.

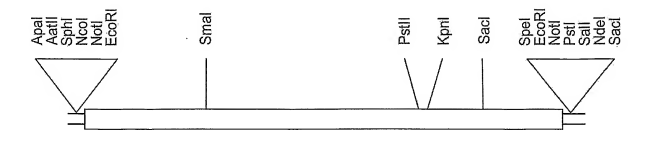
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- 25. A method according to claim 21 wherein said method includes introducing into said plant effective amounts of nucleic acids or nucleic acid fragments encoding all three of CHS or CHS-like, BAN or BAN-like and LAR or LAR-like polypeptides.
- 5 26. Use of a nucleic acid or nucleic acid fragment according to any one of claims 1 to 5, and/or nucleotide sequence information thereof, and/or single nucleotide polymorphisms thereof as a molecular genetic marker.
  - 27. A substantially purified or isolated polypeptide from a clover (*Trifolium*), medic (*Medicago*), ryegrass (*Lolium*) or fescue (*Festuca*) species, selected from the group consisting of CHS and CHS-like, BAN and BAN-like and LAR and LAR-like; and functionally active fragments and variants thereof.

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- 28. A polypeptide according to Claim 27, wherein said polypeptide is from white clover (*Trifolium repens*) or perennial ryegrass (*Lolium perenne*).
- 29. A polypeptide according to Claim 27, wherein said polypeptide is CHS or CHS-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 3, 7, 11 and 15 hereto (Sequence ID Nos. 2, 4, 6 and 8, respectively); and functionally active fragments and variants thereof.
- 30. A polypeptide according to Claim 27, wherein said polypeptide is BAN or BAN-like and includes an amino acid sequence selected from the group consisting of sequence shown in Figure 19 hereto (Sequence ID No. 10); and functionally active fragments and variants thereof.
- 31. A polypeptide according to Claim 27, wherein said polypeptide is LAR or LAR-like and includes an amino acid sequence selected from the group consisting of sequences shown in Figures 23, 27 and 31 hereto (Sequence ID Nos. 12, 14 and 16, respectively); and functionally active fragments and variants thereof.

# 1/40



TrCHSa3

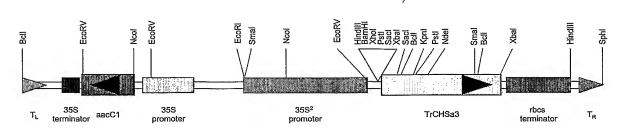
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1	<b>GAATTC</b> ACTA	GTGATTAAGC	AGTGGTAACA	ACGCAGAGTA	CGCGGGGAAC
51	AAAAACAACT	ACGCATATTA	TATATATATA	TATATAGTCT	ATAATTGAAA
101	GAAACTGCTA	AAGATATTAT	TAAGATATGG	TGAGTGTAGC	TGAAATTCGC
151	AAGGCTCAGA	GGGCTGAAGG	CCCTGCAACC	ATTTTGGCCA	TTGGCACTGC
201	AAATCCACCA	AACCGTGTTG	AGCAGAGCAC	ATATCCTGAT	TTCTACTTCA
251	AAATTACAAA	CAGTGAGCAC	AAGACTGAGC	TCAAAGAGAA	GTTCCAACGC
301	ATGTGTGACA	AATCCATGAT	CAAGAGCAGA	TACATGTATC	TAACAGAAGA
351	GATTTTGAAA	GAAAATCCTA	GTCTTTGTGA	ATACATGGCA	CCTTCATTGG
401	ATGCTAGGCA	AGACATGGTG	GTGGTTGAGG	TACCTAGACT	TGGGAAGGAG
451	GCTGCAGTCA	AGGCCATTAA	AGAATGGGGT	CAACCAAAGT	CAAAGATTAC
501	TCACTTAATC	TTTTGCACCA	CAAGTGGTGT	TGACATGCCT	GGTGCTGATT
551	ACCAACTCAC	AAAACTCTTA	GGTCTTCGCC	CATATGTGAA	AAGGTATATG
601	ATGTACCAAC	AAGGTTGTTT	TGCAGGAGGC	ACGGTGCTTC	GTTTGGCAAA
651	AGATTTGGCC	GAGAACAACA	AAGGTGCTCG	TGTGCTAGTT	GTTTGTTCTG
701	AAGTCACCGC	AGTCACATTT	CGCGGCCCCA	GTGATACTCA	CTTGGACAGT
751	${\tt CTTGTTGGAC}$	AAGCATTGTT	TGGAGATGGA	GCCGCTGCAC	TAATTGTTGG
801	TTCTGATCCA	GTGCCTGAAA	TTGAGAAACC	AATATTTGAG	ATGGTTTGGA
851	CTGCACAAAC	AATTGCTCCA	GACAGTGAAG	GTGCCATTGA	TGGTCATCTT
901	CGTGAAGCTG	GGCTAACATT	TCATCTTCTT	AAAGATGTTC	CTGGGATTGT
951	ATCAAAGAAC	ATTAATAAAG	CATTGGTTGA	GGCTTTCCAA	CCATTAGGAA
1001	TTTCTGACTA	CAACTCAATC	TTTTGGATTG	CACACCCGGG	TGGACCTGCA
1051	ATTCTTGATC	AAGTAGAACA	AAAGCTAGCC	TTGAAGCCCG	AAAAGATGAG
1101	GGCCACGAGG	GAAGTTCTAA	GTGAATATGG	AAACATGTCA	AGCGCATGTG
1151	TATTGTTCAT	CTTAGATGAG	ATGCGGAAGA	AATCGGCTCA	AAATGGACTT
1201	AAGACAACTG	GAGAAGGACT	TGATTGGGGT	GTGTTGTTCG	GCTTCGGACC
1251	AGGACTTACC	ATTGAAACCG	TTGTTCTTCG	TAGCGTGGCT	ATATAAGATG
1301	TGTGATTGTT	TTTATTTTAA	TGTATTACTT	TTAATCTTGC	TGCCTTGAAT
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1401	AAAAAAAAA	AAGTACTCTG	CGTTGTTACC	ACTGCTTAAT	CGAATTC

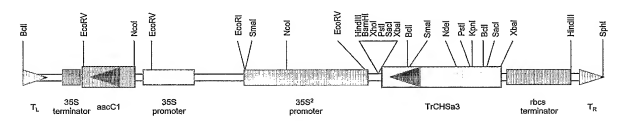
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1	MVSVAEIRKA	QRAEGPATIL	AIGTANPPNR	VEQSTYPDFY	FKITNSEHKT
51	ELKEKFQRMC	DKSMIKSRYM	YLTEEILKEN	PSLCEYMAPS	LDARQDMVVV
101	EVPRLGKEAA	VKAIKEWGQP	KSKITHLIFC	TTSGVDMPGA	DYQLTKLLGL
151	RPYVKRYMMY	QQGCFAGGTV	LRLAKDLAEN	NKGARVLVVC	SEVTAVTFRG
201	PSDTHLDSLV	GQALFGDGAA	ALIVGSDPVP	EIEKPIFEMV	WTAQTIAPDS
251	EGAIDGHLRE	AGLTFHLLKD	VPGIVSKNIN	KALVEAFQPL	GISDYNSIFW
301	IAHPGGPAIL	DQVEQKLALK	PEKMRATREV	LSEYGNMSSA	CVLFILDEMR
351	KKSAONGLKT	TGEGLDWGVL	FGFGPGLTIE	TVVLRSVAI	

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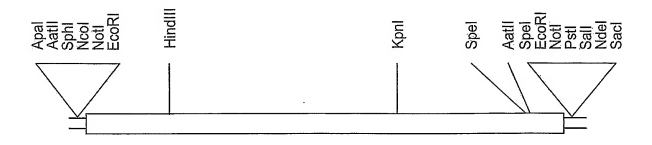


pPZP221:35S2TrCHSa3 sense



pPZP221:35S2TrCHSa3 anti

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**TrCHSc** 

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			0, . 0		
1	$\mathbf{GAATTC}\mathbf{GATT}$	AAGCAGTGGT	AACAACGCAG	AGTACGCGGG	GATTCAATCT
51	GTTGTGCATA	AAATTCACTC	ATTGCATAGA	AAACCATACA	CATTTGATCT
101	TGCAAAGAAG	AAATATGGGA	GACGAAGGTA	TAGTGAGAGG	TGTCACAAAG
151	CAGACAACCC	$\tt CTGGGAAGGC$	TACTATATTG	$\tt GCTCTTGGCA$	AGGCATTCCC
201	TCACCAACTT	GTGATGCAAG	AGTGTTTAGT	${\tt TGATGGTTAT}$	TTTAGGGACA
251	CTAATTGTGA	CAATCCTGAA	CTTAAGCAGA	AACTTGCTAG	ACTTTGTAAG
301	ACAACCACGG	TAAAAACAAG	GTATGTTGTT	ATGAATGAGG	AGATACTAAA
351	GAAATATCCA	GAACTTGTTG	TCGAAGGCGC	CTCAACTGTA	AAACAACGTT
401	TAGAGATATG	TAATGAGGCA	GTAACACAAA	TGGCAATTGA	AGCTTCCCAA
451	GTTTGCCTAA	AGAATTGGGG	TAGATCCTTA	TCGGACATAA	CTCATGTGGT
501	TTATGTTTCA	TCTAGTGAAG	CTAGATTACC	${\tt CGGTGGTGAC}$	CTATACTTGT
551	CAAAAGGACT	AGGACTAAAC	CCTAAAATTC	AAAGAACCAT	GCTCTATTTC
601	TCTGGATGCT	CGGGAGGCGT	AGCCGGCCTT	${\tt CGCGTTGCGA}$	AAGACGTAGC
651	TGAGAACAAC	CCTGGAAGTA	GAGTTTTGCT	TGCTACTTCG	GAAACTACAA
701	${\tt TTATTG}_{\!$	CAAGCCACCA	AGTGTTGATA	GACCTTATGA	TCTTGTTGGT
751	GTGGCACTCT	TTGGAGATGG	TGCTGGTGCA	ATGATAATTG	GCTCAGACCC
801	GGTATTTGAA	ACTGAGACAC	CATTGTTTGA	GCTGCATACT	TCAGCTCAGG
851	AGTTTATACC	AGACACCGAG	AAGAAAATTG	ATGGGCGGCT	GACGGAGGAG
901	GGCATAAGTT	TCACACTAGC	AAGGGAACTT	CCGCAGATAA	TCGAAGACAA
951	TGTTGAGGGA	TTCTGTAATA	AACTAATTGA	TGTTGTTGGG	TTGGAGAATA
1001	AGGAGTACAA	TAAGTTGTTT	TGGGCTGTGC	ATCCAGGTGG	GCCTGCGATA
1051	TTGAATCGCG	TGGAGAAGCG	GCTTGAGTTG	TCGCCGCAGA	AGCTGAATGC
1101	TAGTAGAAAA	GCTCTAATGG	ATTATGGAAA	TGCTAGCAGC	AATACTATTG
1151	TTTATGTGCT	GGAATATATG	CTAGAAGAGG	AAAAGAAGAT	TAAAAAGGCG
1201	GGTGGAGGAG	ATTCTGAATG	GGGATTGATA	CTTGCTTTTG	GACCTGGAAT
1251	TACTTTTGAG	GGGATTCTAG	CAAGGAACTT	GTGTGCATGA	AGTCTTATAC
1301	AATTGTGATG	CATGACTTAT	ACTCTTATTT	CTACTAATTA	TTATATTAAG
1351	CAAATTCAGA	ACTTTTAAGT	AATGATTTAA	TGAAGAATAC	TTATAGTATA
1401	TTGACTTTAT	TCACTTTCAA	AGCAAGTTTA	TGATCCTAAG	ACATGGTAGA
1451	ACTTGAGCAT	GTGGAATAGT	TGTAACAAAA	ACTCTAAGCA	AATAGAGACT
1501	TTATGTAGTA	TAAAGCATTT	CCAGACATGA	TAAATAATGG	TACCTCAGAA
1551	CATAAAATAT	ATTTAGCTAT	CTTTCATCCC	CAACTTTACA	CATCCACCAA
1601	GGTACAGAAT	AAGCATATGT	CAACACAAAA	TGTACTCTAA	GTCTAACATG
1651	AGTAACCAAA	CATGATGCCT	GATTAAGTTA	AAAGAAAAGA	AAATCTGAGG
1701	GCATAGATCT	TCAATCACAC	CACTCCAGAG	GGAAGGCGTA	GAACAAGCTG

#### 7/40

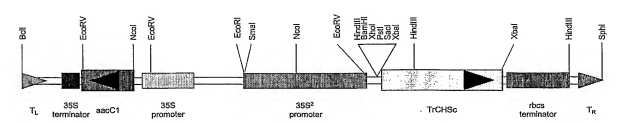
. / 5 1	ICCGCCGAAA	ACACIGCAAI	ICAAIAAAIA	ICALIAGGAC	DAJDIDAJAA
.801	AGTCATGCGG	GAAATGTCTT	AAGTCACTGT	ACTAAAAATA	TAGGATTATA
851	TTATGAACTA	TACTAACCTT	TTCACATAAT	AGTAACAGAA	ATCAGCTAAG
901	ATGAATGTCT	GGACAATTTC	TGAGATAAGA	ACCATGACGG	CCATAAGCCA
951	TACCCCAAGG	CAACCAATAA	ATGTCCACGG	GTATCTAACA	CCTGTTGCAA
2001	GAAATAGTAA	GTTATTAGGA	GATGTGCGGT	TACGAAATTC	AAGCTACACA
2051	ACAAAAGGAG	GCCAGAACAA	CAGCAATCTT	GTAACCAGAT	GACAACAATA
2101	AAATGTAAAC	TTAAAGAGAC	CGAACACACA	AACATTGCAA	CTCAGATGGA
2151	ATTGCTGCCA	TGTAACTAGT	AGGAGATTTG	GGACGTCAAA	TCAGTATATT
2201	ATGCAAATAC	AAGGTATGAC	CGCCTTGTCT	ATTGTAGCAT	ACAACAAACG
2251	TACAGTGGGT	TTGTCCCTCT	CAAAATGGCA	${\tt GGATCTTTAC}$	AGCACAATAT
2301	TTGGTTTTGT	CATACTTATA	CCATAAAAAA	AAAAAAAAA	ААААААААА
2351	AAAGTACTCT	GCGTTGTTAC	CACTGCTTAA	TCACTAGT <b>G</b> A	ATTC

# FIGURE 6 (cont.)

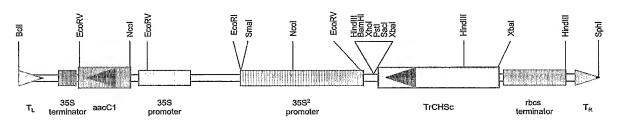
#### 8/40

Т.	MGDEGTAKGA	INGITEGRAL	Thangvarbu	QT AMORCTAD	GYFRDINCDN
51	PELKQKLARL	CKTTTVKTRY	VVMNEEILKK	YPELVVEGAS	TVKQRLEICN
101	EAVTQMAIEA	SQVCLKNWGR	SLSDITHVVY	VSSSEARLPG	GDLYLSKGLG
151	LNPKIQRTML	YFSGCSGGVA	GLRVAKDVAE	NNPGSRVLLA	TSETTIIGFK
201	PPSVDRPYDL	VGVALFGDGA	GAMIIGSDPV	FETETPLFEL	HTSAQEFIPD
251	TEKKIDGRLT	EEGISFTLAR	ELPQIIEDNV	EGFCNKLIDV	VGLENKEYNK
301	LFWAVHPGGP	AILNRVEKRL	ELSPQKLNAS	RKALMDYGNA	SSNTIVYVLE
351	YMLEEEKKIK	KAGGGDSEWG	LILAFGPGIT	FEGILARNLC	A

#### 9/40

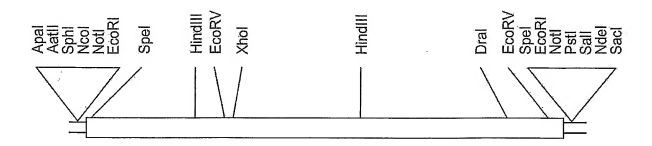


pPZP221:35S2TrCHSc sense



pPZP221:35S2TrCHSc anti

## 10/40



**TrCHSf** 

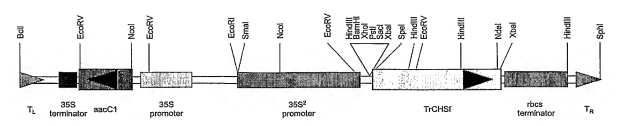
## 11/40

1	GAATTC GATT	AAGCAGTGGT	AACAACGCAG	AGTACGCGGG	ACTAAGCCTT
51	GATTCATTGT	TTGTTTCCAT	AACACAAGAA	CTAGTGTTTG	CTTGAATCTT
101	AAGAAAAAT	GCCTCAAGGT	GATTTGAATG	GAAGTTCCTC	GGTGAATGGA
151	GCACGTGCTA	GACGTGCTCC	TACTCAGGGA	AAGGCAACGA	TACTTGCATT
201	AGGAAAGGCT	TTCCCCGCCC	AGGTCCTCCC	TCAAGAGTGC	TTGGTGGAAG
251	GATTCATTCG	CGACACTAAG	TGTGACGATA	CTTATATTAA	GGAGAAATTG
301	GAGCGTCTTT	GCAAAAACAC	AACTGTGAAA	ACAAGATACA	CAGTAATGTC
351	AAAGGAGATC	TTAGACAACT	ATCCAGAGCT	AGCCATAGAT	GGAACACCAA
401	CAATAAGGCA	AAAGCTTGAA	ATAGCAAATC	CAGCAGTAGT	TGAAATGGCA
451	ACAAGAGCAA	GCAAAGATTG	CATCAAAGAA	TGGGGAAGGT	CACCTCAAGA
501	TATCACACAC	ATAGTCTATG	TTTCCTCGAG	CGAAATTCGT	CTACCCGGTG
551	GTGACCTTTA	TCTTGCAAAT	GAACTCGGCT	TAAACAGCGA	TGTTAATCGC
601	GTAATGCTCT	ATTTCCTCGG	TTGCTACGGC	GGTGTCACTG	GCTTACGTGT
651	CGCCAAAGAC	ATCGCCGAAA	ATAACCCTGG	TAGTAGGGTG	TTACTCACAA
701	CATCCGAGAC	CACTATTCTC	GGTTTTCGAC	CACCGAGTAA	AGCTAGACCT
751	TATGACCTCG	TTGGCGCTGC	ACTTTTCGGT	GATGGCGCCG	CTGCTGCAAT
801	AATTGGAACA	GACCCTATAT	TGAATCAAGA	ATCACCTTTC	ATGGAATTGA
851	ACCATGCAGT	CCAAAAATTC	TTGCCTGATA	CACAAAATGT	GATTGATGGT
901	AGAATCACTG	AAGAGGGTAT	${\tt TAATTTTAAG}$	${\tt CTTGGAAGAG}$	ACCTTCCTCA
951	AAAAATTGAA	GACAATATTG	AAGAATTTTG	CAAGAAAATT	ATGGCTAAAA
1001	GTGATGTTAA	GGAATTTAAT	GACTTATTTT	$\tt GGGCTGTTCA$	TCCTGGTGGG
1051	CCAGCTATAC	TCAATAAGCT	AGAAAATATA	CTCAAATTGA	AAAGTGATAA
1101	ATTGGATTGT	AGTAGGAAGG	CATTAATGGA	TTATGGAAAT	GTTAGTAGCA
1151	ATACTATATT	CTATGTGATG	GAGTATATGA	GAGATTATTT	GAAGGAAGAT
1201	GGAAGTGAAG	AATGGGGATT	AGGATTGGCT	TTTGGACCAG	GGATTACTTT
1251	TGAAGGGGTT	CTCCTCCGTA	GCCTTTAATC	TTGAAATAAT	AATTCATATG
1301	AAATTACTTG	TCTTAAGATT	GTGATAGGAA	GATGAATATG	TATTGGATTA
1351	ATATTGATAT	GGTGTTATTT	TAAGTTGATT	TTAAAAAAAG	TTTATTAATA
1401	AAGTATGATG	TAACAATTGT	TGTTTGAATG	TTAAAAGGGA	AGTATACTAT
1451	TTTAAGTTCT	TGACCATACT	GATTTTTTCT	TTACACATTT	TCATATCTAA
1501	AATTGTTCTA	TGATATCTTC	ATTGTTGATA	CTGTAATAAT	ATAATATCTA
1551	ATTTGGCTGG	CAAAATGAAA	GATTTTTCAC	CGAAAAAAAA	AAAAAAAA
1601	AAAAAAAAA	AAGTACTCTG	CGTTGTTACC	ACTGCTTAAT	CACTAGTGAA
1651	TTC				

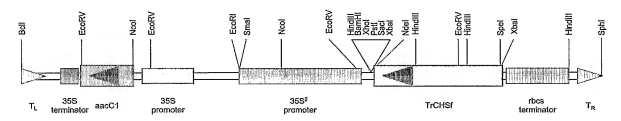
## 12/40

1	MPQGDLNGSS	SVNGARARRA	PTQGKATILA	LGKAF. PAQVL	POECTAEGET
51	RDTKCDDTYI	KEKLERLCKN	TTVKTRYTVM	SKEILDNYPE	LAIDGTPTIR
101	QKLEIANPAV	VEMATRASKD	CIKEWGRSPQ	DITHIVYVSS	SEIRLPGGDL
151	YLANELGLNS	DVNRVMLYFL	GCYGGVTGLR	VAKDIAENNP	GSRVLLTTSE
201	TTILGFRPPS	KARPYDLVGA	ALFGDGAAAA	IIGTDPILNQ	ESPFMELNHA
251	VQKFLPDTQN	VIDGRITEEG	INFKLGRDLP	QKIEDNIEEF	CKKIMAKSDV
301	KEFNDLFWAV	HPGGPAILNK	LENILKLKSD	KLDCSRKALM	DYGNVSSNTI
351	FYVMEYMRDY	LKEDGSEEWG	LGLAFGPGIT	FEGVLLRSL	

#### 13/40

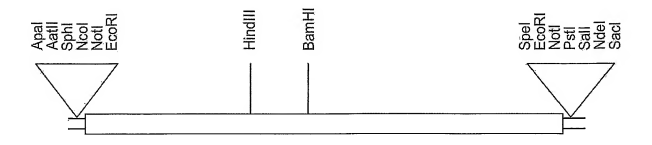


pPZP221:35S2TrCHSf sense



pPZP221:35S2TrCHSf anti

## 14/40



**TrCHSh** 

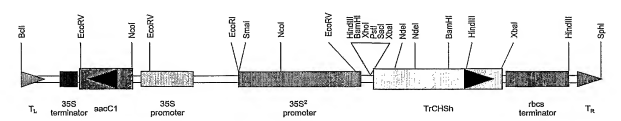
# 15/40

1	<b>GAATTC</b> ACTA	GTGATTAAGC	AGTGGTAACA	ACGCAGAGTA	CGCGGGGGAA
51	TCCACCAAAT	CAACACCATT	AATAACCTTC	CAAATTCTCG	TTACCTCACC
101	AAATCTCATT	TTTCATTATA	TATCTTGGGT	ACATCTTTTG	TTACCTCCAA
151	CAAAAAAATG	GTGACCGTAG	AAGAGATTCG	TAACGCCCAA	CGTTCAAATG
201	GCCCTGCCAC	TATCTTAGCT	TTTGGCACAG	CCACTCCTTC	TAACTGTGTC
251	ACTCAAGCTG	ATTATCCTGA	TTACTACTTT	CGTATCACCA	ACAGCGAACA
301	TATGACTGAT	CTTAAGGAAA	AATTCAAGCG	GATGTGTGAT	AGATCAATGA
351	TAAAGAAACG	TTACATGCAC	CTAACAGAAG	ACTTTCTGAA	GGAGAATCCA
401	AATATGTGTG	AATACATGGC	ACCATCACTA	GATGTAAGAC	GAGACATAGT
451	GGTTGTTGAA	GTACCAAAGC	TAGGTAAAGA	AGCAGCAAAA	AAAGCCATAT
501	${\tt GTGAATGGGG}$	ACAACCAAAA	TCCAAAATCA	CACATCTTGT	TTTCTGCACC
551	ACTTCCGGTG	TTGACATGCC	GGGAGCCGAT	TACCAACTCA	CCAAACTTTT
601	AGGCTTAAAA	CCTTCTGTCA	AGCGTCTCAT	GATGTATCAA	CAAGGTTGTT
651	TCGCTGGCGG	CACAGTTCTC	CGCTTAGCAA	AAGACCTTGT	TGAGAATAAC
701	AAAAATGCAA	GAGTTCTTGT	TGTTTGTTCT	GAAATTACTG	CGGTTACTTT
751	TCGTGGACCA	TCGGATACTC	ATCTTGATTC	$\tt GCTCGTGGGA$	CAGGCGCTTT
801	TTGGTGATGG	AGCCGCAGCA	ATGATTATTG	GTGCGGATCC	TGATTTAACC
851	GTGGAGCGTC	${\tt CGATTTTCGA}$	GATTGTTTCG	GCTGCTCAGA	CTATTCTTCC
901	TGATTCTGAT	GGCGCAATTG	ATGGACATCT	TCGTGAAGTG	GGGCTCACTT
951	TTCATTTATT	GAAAGATGTT	CCGGGGATTA	TTTCAAAGAA	CATTGAAAAA
1001	AGTTTAGTTG	AAGCTTTTGC	GCCTATTGGG	ATTAATGATT	GGAACTCAAT
1051	ATTTTGGGTT	GCACATCCAG	GTGGACCGGC	TATTTTAGAC	CAGGTTGAAG
1101	AGAAACTCCA	TCTTAAAGAG	GAGAAACTCC	GGTCCACCCG	GCATGTGCTT
1151	AGTGAATATG	GAAATATGTC	AAGTGCATGT	$\tt GTTTTATTTA$	TTTTGGATGA
1201	AATGAGAAAG	AGGTCTAAAG	AGGAAGGGAT	GATTACAACT	GGTGAAGGGT
1251	TGGAATGGGG	TGTGTTGTTT	GGGTTTGGAC	CGGGTTTAAC	TGTTGAAACC
1301	GTTGTGCTTC	ATAGTGTTCC	GGTTCAGGGT	TGAATTTATT	ATACATAGAT
1351	TGGAAAATAA	AATTTGCCTG	CCGAGAGATG	TGAACTAACT	TTGTAGGCAA
1401	GCTCAAATTA	AAGTTTGAGA	TAATATTGTG	CTTTAGTTAT	TATGGTATGT
1451	AATGTAATGT	TTTTACTTTT	TTCGAAATTC	ATGTAATTTG	ATATGTAAAG
1501	TAATATGTTT	GGGTTGGAAT	ATAATTATTT	GTTAACTAAA	AAAAAAAAA
1551	AAAAAAAAA	AAAAAGTACT	CTGCGTTGTT	ACCACTGCTT	AATC <b>GAATTC</b>

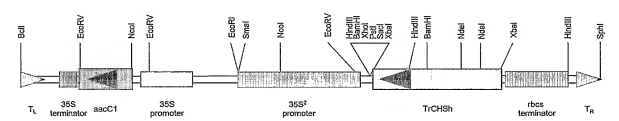
## 16/40

1.	MVTVEEIRNA	QRSNGPATIL	AFGTATPSNC	VTQADYPDYY	FRITNSEHMT
51	DLKEKFKRMC	DRSMIKKRYM	HLTEDFLKEN	PNMCEYMAPS	LDVRRDIVVV
101	EVPKLGKEAA	KKAICEWGQP	KSKITHLVFC	TTSGVDMPGA	DYQLTKLLGI
151	KPSVKRLMMY	QQGCFAGGTV	LRLAKDLVEN	NKNARVLVVC	SEITAVTFRO
201	PSDTHLDSLV	GQALFGDGAA	AMIIGADPDL	TVERPIFEIV	SAAQTILPDS
251	DGAIDGHLRE	VGLTFHLLKD	VPGIISKNIE	KSLVEAFAPI	GINDWNSIFW
301	VAHPGGPAIL	DQVEEKLHLK	EEKLRSTRHV	LSEYGNMSSA	CVLFILDEME
351	KRSKEEGMIT	TGEGLEWGVL	FGFGPGLTVE	TVVLHSVPVQ	G

#### 17/40

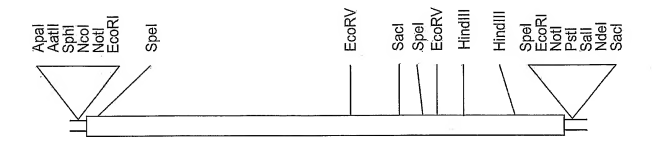


pPZP221:35S2TrCHSh sense



pPZP221:35S2TrCHSh anti

#### 18/40



**TrBANa** 

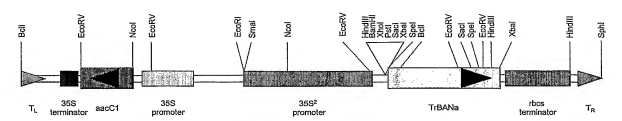
#### 19/40

1	<b>GAATTC</b> GATT	AAGCAGTGGT	AACAACGCAG	AGTACGCGGG	ATAAAAACTG
51	CACTAGTGTG	TATAAGTTTC	TTGGTGAAAA	AAGAGTTTGT	AAATTAACAT
101	CATGGCTAGT	ATCAAACAAA	TTGGAAACAA	GAAAGCATGT	GTGATTGGTG
151	GCACTGGTTT	TGTTGCATCT	ATGTTGATCA	AGCAGTTACT	TGAAAAGGGT
201	TATGCTGTTA	ATACTACCGT	TAGAGACCCA	GATAGCCCTA	AGAAAATATC
251	TCACCTAGTG	GCACTGCAAA	$\tt GTTTGGGGGA$	ACTGAATCTA	TTTAGAGCAG
301	ACTTAACAGT	TGAAGAAGAT	TTTGATGCTC	CTATAGCAGG	ATGTGAACTT
351	GTTTTTCAAC	TTGCTACACC	TGTGAACTTT	GCTTCTCAAG	ATCCTGAGAA
401	TGACATGATA	AAGCCAGCAA	TCAAAGGTGT	GTTGAATGTG	TTGAAAGCAA
451	TTGCAAGAGC	AAAAGAAGTT	AAAAGAGTTA	TCTTAACATC	TTCGGCAGCC
501	GCGGTGACTA	TAAATGAACT	CAAAGGGACA	GGTCATGTTA	TGGATGAAAC
551	CAACTGGTCT	GATGTTGAAT	TTCTCAACAC	TGCAAAACCA	CCCACTTGGG
601	GTTATCCTGC	CTCAAAAATG	CTAGCTGAAA	AGGCTGCATG	GAAATTTGCT
651	GAAGAAAATG	ACATTGATCT	AATCACTGTG	ATACCTAGTT	TAACAACTGG
701	TCCTTCTCTC	ACACCAGATA	TCCCATCTAG	TGTTGGCTTG	GCAATGTCTC
751	TAATAACAGG	CAATGATTTT	CTCATAAATG	CTTTGAAAGG	AATGCAGTTT
801	CTGTCGGGTT	CGTTATCCAT	CACTCATGTT	GAGGATATTT	GCCGAGCTCA
851	TATATTTCTT	GCAGAGAAAG	AATCAGCTTC	TGGTAGATAC	ATTTGCTGTG
901	CTCACAATAC	TAGTGTTCCC	GAGCTTGCAA	AGTTTCTCAA	CAAACGATAT
951	CCTCAGTATA	AAGTTCCAAC	TGAATTTGAT	GATTGCCCCA	GCAAGGCAAA
1001	GTTGATAATC	TCTTCTGAAA	AGCTTATCAA	AGAAGGGTTC	AGTTTCAAGC
1051	ATGGTATTGC	CGAAACTTTC	GACCAGACTG	TCGAGTATTT	TAAGACTAAG
1101	GGGGCACTGA	AGAATTAGAT	TTTGATATTT	CTAATTCAAT	AGCAAACTCT
1151	AAGCTTGTTA	TGTGTTTGTG	AAGTTCAGAG	TGAAATATCA	AATGAATAAG
1201	TGGAGAGAGC	ACAATAAGAG	GAGAGCACAA	TAATTTTGGA	AAAAAAAAA
1251	AAAAAAAAA	AAAAAAAAGT	ACTCTGCGTT	GTTACCACTG	CTTAATCACT
1301	AGT <b>GAATTC</b>				

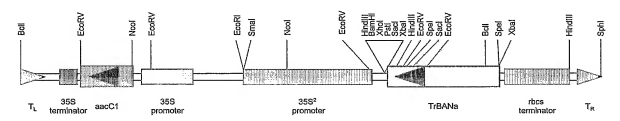
## 20/40

1	MASIKQIGNK	KACVIGGTGF	VASMLIKQLL	EKGYAVNTTV	RDPDSPKKIS
51	HLVALQSLGE	LNLFRADLTV	EEDFDAPIAG	CELVFQLATP	VNFASQDPEN
101	DMIKPAIKGV	LNVLKAIARA	KEVKRVILTS	SAAAVTINEL	KGTGHVMDET
151	NWSDVEFLNT	AKPPTWGYPA	SKMLAEKAAW	KFAEENDIDL	ITVIPSLTTO
201	PSLTPDIPSS	VGLAMSLITG	NDFLINALKG	MQFLSGSLSI	THVEDICRAF
251	IFLAEKESAS	GRYICCAHNT	SVPELAKFLN	KRYPQYKVPT	EFDDCPSKA
301	LIISSEKLIK	EGFSFKHGIA	ETFDOTVEYF	KTKGALKN	

#### 21/40

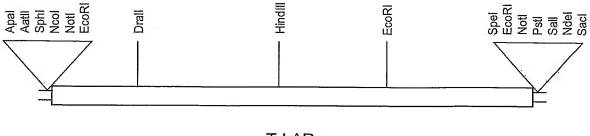


pPZP221:35S2TrBANa sense



pPZP221:35S2TrBANa anti

#### 22/40



TrLARa

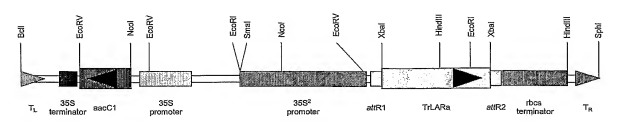
#### 23/40

1	GAATTCGATT	AAGCAGTGGT	AACAACGCAG	AGTACGCGGG	GATACCAACA
51	TTGTCACAAT	TAACTCTAAA	AGCAAAGCAA	TGGCACCAGC	AGCAACATCA
101	TCACCAACCA	CTCCTACTAC	TACCAAGGGT	CGTGTCCTAA	TTGTTGGAGG
151	AACAGGTTTC	ATTGGAAAAT	TTGTAACTGA	GGCAAGTCTT	TCCACAACAC
201	ACCCAACCTA	CTTGTTGGTT	CGGCCAGGAC	CTCTTCTCTC	TTCTAAGGCT
251	GCCACTATTA	AGGCATTCCA	AGAGAAAGGT	GCCATTGTCA	TTTATGGTCG
301	GGTAAATAAT	AAGGAGTTCA	TGGAGATGAT	TTTGAAAAAG	TATGAGATAA
351	ATGTAGTCAT	TTCTGCAATA	GGAGGCTCTG	ATGGCTTGCT	GGAACAGCTT
401	ACTTTGGTGG	AGGCCATGAA	ATCTATTAAC	ACCATTAAGA	GGTTTTTGCC
451	TTCGGAATTT	GGTCACGATG	TGGACAGAGC	AAATCCTGTG	GAACCTGGCC
501	TAACAATGTA	CAAACAGAAA	${\tt CGTTTGGTTA}$	GACGTGTGAT	CGAAGAATCT
551	GGTATACCAT	ACACCTACAT	CTGTTGCAAT	TCGATCGCAT	CTTGGCCGTA
601	CTATGACAAT	TGTCATCCAT	CACAGCTTCC	TCCACCGTTG	GATCAATTAC
651	ATATTTATGG	TCATGGCGAT	GTCAAAGCTT	ACTTTGTTGA	TGGCTATGAT
701	ATTGGGAAAT	TCACAATGAA	GGTCATTGAT	GATGAAAGAA	CAATCAACAA
751	AAATGTTCAT	TTTCGACCTT	CTAACAATTG	TTATAGCATG	AATGAGCTTG
801	CTTCTTTGTG	GGAAAACAAA	ATTGCACGAA	AAATTCCTAG	AGTGATCGTC
851	TCTGAAGACG	ATCTTCTAGC	AATAGCCGCA	GAAAATTGCA	TACCGGAAAG
901	TGTCGTGGCA	CCAATCACTC	ATGATATATT	CATCAATGGA	TGTCAAGTTA
951	ACTTCAAGAT	AGATGGAATT	CATGATGTTG	AAATTGGCAC	TCTATATCCT
1001	GGTGAATCGG	TAAGAAGTTT	GGAGGAATGC	TATGAGAAAT	TTGTTGTCAT
1051	GGCGGCTGAC	AAGATTCATA	AAGAAGAAAC	TGGAGTTACC	GCAGGTGGGG
1101	GCGGCACAAC	GGCTATGGTA	GAGCCGGTGC	CAATCACAGC	TTCCTGTTGA
1151	AAAGGTTCAC	CTGAGGTGGA	TATTCTTTTG	AGTCATAAGA	CATGTTGATT
1201	GTTGATGTTG	TTTTCAAGAA	TGTTTCATCA	TTTCATGTGT	TTTATTAATC
1251	CTAAGTACAA	ATAATTGCTG	TCTACGTACG	TTCTTAGTTG	CAAAAATTCT
1301	TGTTATTCTC	TATTGAGGTA	AAAGTCTTCA	TGTTTACAAA	AAAAAAAAA
1351	AAAAAAAAA	AAAAAAAAGT	ACTCTGCGTT	GTTACCACTG	CTTAATCACT
1401	AGTGAATTC				

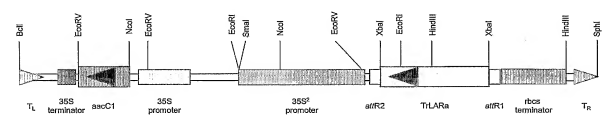
#### 24/40

1	MAPAATSSPT	TPTTTKGRVL	IVGGTGFIGK	FVTEASLSTT	HPTYLLVRPO
51	PLLSSKAATI	KAFQEKGAIV	IYGRVNNKEF	MEMILKKYEI	NVVISAIGGS
101	DGLLEQLTLV	EAMKSINTIK	RFLPSEFGHD	VDRANPVEPG	LTMYKQKRLV
151	RRVIEESGIP	YTYICCNSIA	SWPYYDNCHP	SQLPPPLDQL	HIYGHGDVKA
201	YFVDGYDIGK	FTMKVIDDER	TINKNVHFRP	SNNCYSMNEL	ASLWENKIAF
251	KIPRVIVSED	DLLAIAAENC	IPESVVAPIT	HDIFINGCQV	NFKIDGIHDV
301	EIGTLYPGES	VRSLEECYEK	FVVMAADKIH	KEETGVTAGG	GGTTAMVEPV
351	PITASC				

#### 25/40



pPZP221:35S2TrLARa sense



pPZP221:35S2TrLARa anti

## 26/40

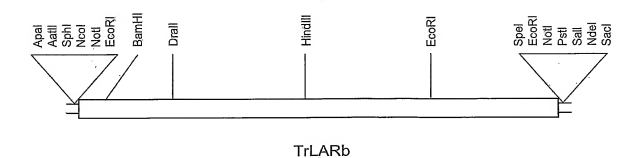


FIGURE 25

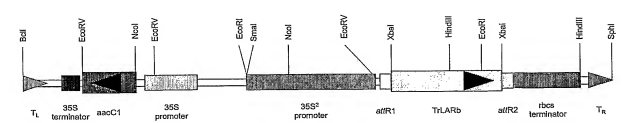
#### 27/40

1	<b>GAATTC</b> GATT	AAGCAGTGGT	AACAACGCAG	AGTACGCGGG	AGGATCCTTC
51	CATTTTGCAT	ACCAACATTG	TCACAATTAA	CTCTAAAAGC	AAAGCAATGG
101	CACCAGCAGC	AACATCATCA	CCAACCACTC	CTACTACTAC	CAAGGGTCGT
151	GTCCTAATTG	TTGGAGGAAC	AGGTTTCATT	GGAAAATTTG	TAACTGAGGC
201	AAGTCTTTCC	ACAACACACC	CAACCTACTT	GTTGGTTCGG	CCAGGACCTC
251	TTCTCTCTTC	TAAGGCTGCC	ACTATTAAGG	CATTCCAAGA	GAAAGGTGCC
301	ATTGTCATTT	ATGGTCGGGT	AAATAATAAG	GAGTTCATGG	AGATGATTTT
351	GAAAAAGTAT	GAGATAAATG	TAGTCATTTC	TGCAATAGGA	GGCTCTGATG
401	GCTTGCTGGA	ACAGCTTACT	TTGGTGGAGG	CCATGAAATC	TATTAACACC
451	ATTAAGAGGT	TTTTGCCTTC	AGAATTTGGT	CACGATGTGG	ACAGAGCAAA
501	TCCTGTGGAA	CCTGGCCTAA	CAATGTACAA	ACAGAAACGT	TTGGTTAGAC
551	GTGTGATCGA	AGAATCTGGT	GTACCATACA	CCTACATCTG	TTGCAATTCG
601	ATCGCATCCT	GGCCGTACTA	TGACAATTGT	CATCCATCAC	AGCTTCCTCC
651	ACCGTTGGAT	CAATTACATA	TTTATGGTCA	TGGCGATGTC	AAAGCTTACT
701	TTGTTGATGG	CTATGATATT	GGGAAATTCA	CAATGAAGGT	CATTGATGAT
751	GAAAGAACAA	TCAACAAAAA	TGTTCATTTT	CGACCTTCTA	ACAATTGTTA
801	TAGCATGAAT	GAGCTTGCTT	CTTTGTGGGA	AAACAAAATT	GCACGAAAAA
851	TTCCTAGAGT	GATCGTCTCT	GAAGACGATC	TTCTAGCAAT	AGCCGCAGAA
901	AACTGCATAC	CGGAAAGTGT	TGTGGCATCA	ATCACTCATG	ATATATTCAT
951	CAATGGATGT	CAAGTTAACT	TCAAGGTAGA	TGGAATTCAT	GATGTTGAAA
1001	TTGGCACTCT	ATATCCTGGT	GAATCGGTAA	GAAGTTTGGA	GGAATGCTAT
1051	GAGAAATTTG	TTGTCATGGC	GGCTGACAAG	ATTCATAAAG	AAGAAACTGG
1101	AGTTACCGCA	GGTGGGGGCG	GCACAACGGC	TATGGTAGAG	CCGGTGCCAA
1151	TCACAGCTTC	CTGTTGAAAA	GGTTCACCTG	AGGTGGATAT	TCTTTTGAGT
1201	CATAAGACAT	GTTGATTGTT	GATGTTGTTT	TCAAGAATGT	TTCATCATTT
1251	CATGTGTTTT	ATTAATCCTA	AGTACAAATA	ATTGCTGTCT	ACGTACGTTC
1301	TTAGTTGCGA	AAATTCTTGT	TATTCTCTAT	TGGGGTAAAA	GTCTTCATGT
1351	TTATTGTAGT	TGTGTTGGTT	TTTCATATAT	GCTATTTGCA	ATAATGATTT
1401	TTGTGAAGCA	CTTGTGGTGT	ATTTACTTAC	TACTGAAAAT	AATGGTTACA
1451	CAAAATATAT	AAAAAAATAA	AAATAAGCAA	AAAAAAAAA	AAAAAAAAA
1501	AAAAAAAA	GTACTCTGCG	TTGTTACCAC	TGCTTAATCA	CTAGT <b>GAATT</b>
1551	C				

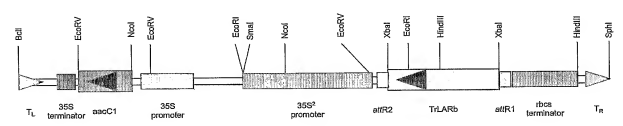
#### 28/40

1	MAPAATSSPT	TPTTKGRVL	IVGGTGFTGK	FVTEASLSTT	HELITTAKEG
51	PLLSSKAATI	KAFQEKGAIV	IYGRVNNKEF	MEMILKKYEI	NVVISAIGGS
101	DGLLEQLTLV	EAMKSINTIK	RFLPSEFGHD	VDRANPVEPG	LTMYKQKRLV
151	RRVIEESGVP	YTYICCNSIA	SWPYYDNCHP	SQLPPPLDQL	HIYGHGDVKA
201	YFVDGYDIGK	FTMKVIDDER	TINKNVHFRP	SNNCYSMNEL	ASLWENKIAF
251	KIPRVIVSED	DLLAIAAENC	IPESVVASIT	HDIFINGCQV	NFKVDGIHDV
301	EIGTLYPGES	VRSLEECYEK	FVVMAADKIH	KEETGVTAGG	GGTTAMVEPV
351	PITASC				

#### 29/40

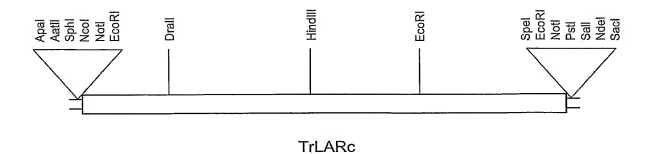


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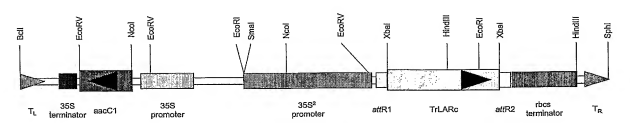
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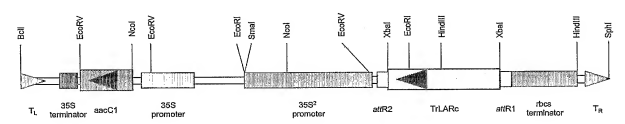
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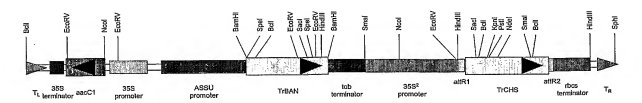


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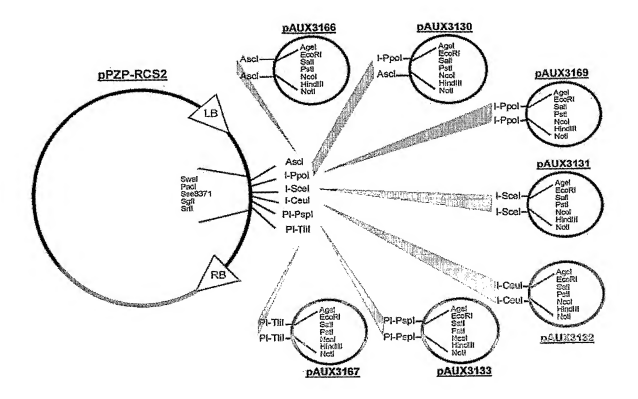
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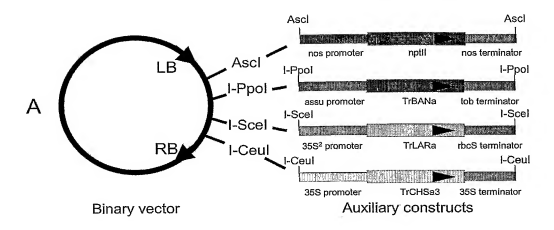


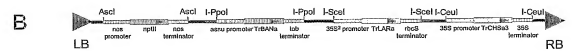
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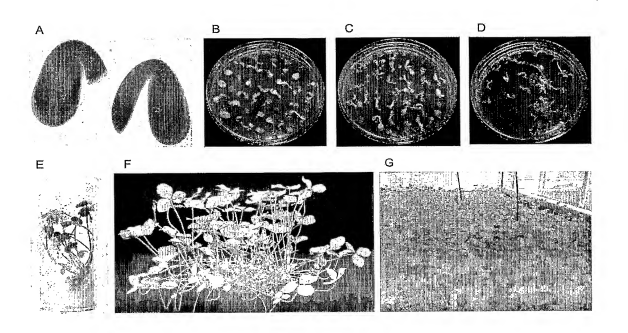
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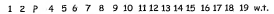


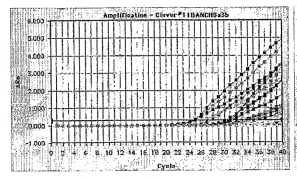
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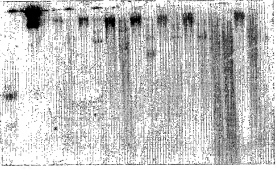
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## 38/40

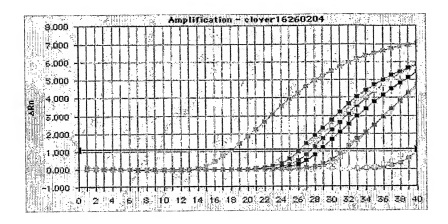




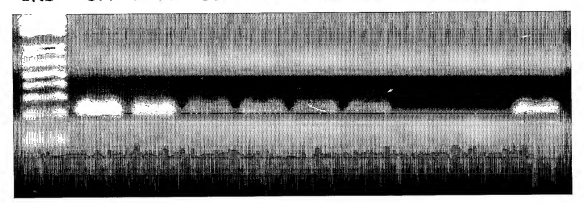




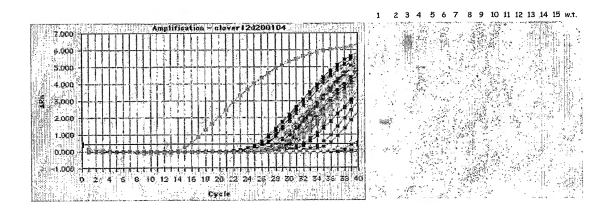
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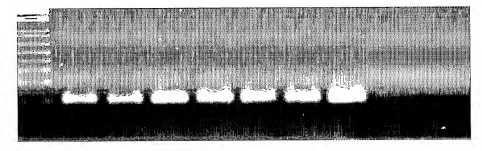
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Met Val Val Glu Val Pro Arg Leu Gly Lys Glu Ala Ala Val Lys 100 105 110

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Pro Leu Gly Ile Ser Asp Tyr Asn Ser Ile Phe Trp Ile Ala His Pro 290 295 300

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Asp Asn Pro Glu Leu Lys Gln Lys Leu Ala Arg Leu Cys Lys Thr Thr 50 60

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Tyr Pro Glu Leu Val Val Glu Gly Ala Ser Thr Val Lys Gln Arg Leu 85 90 95

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Glu Arg Leu Cys Lys Asn Thr Thr Val Lys Thr Arg Tyr Thr Val Met 65 70 75 80

Ser Lys Glu Ile Leu Asp Asn Tyr Pro Glu Leu Ala Ile Asp Gly Thr 85 90 95

Pro Thr Ile Arg Gln Lys Leu Glu Ile Ala Asn Pro Ala Val Val Glu 100 105 110

Met Ala Thr Arg Ala Ser Lys Asp Cys Ile Lys Glu Trp Gly Arg Ser Page 7 M80676490.ST25 115 120 12

Pro Gln Asp Ile Thr His Ile Val Tyr Val Ser Ser Ser Glu Ile Arg 130 135 140 Leu Pro Gly Gly Asp Leu Tyr Leu Ala Asn Glu Leu Gly Leu Asn Ser 145 150 155 160 Asp Val Asn Arg Val Met Leu Tyr Phe Leu Gly Cys Tyr Gly Gly Val 165 170 175 Thr Gly Leu Arg Val Ala Lys Asp Ile Ala Glu Asn Asn Pro Gly Ser 180 185 190 Arg Val Leu Leu Thr Thr Ser Glu Thr Thr Ile Leu Gly Phe Arg Pro 195 200 205 Pro Ser Lys Ala Arg Pro Tyr Asp Leu Val Gly Ala Ala Leu Phe Gly 210 215 220 Asp Gly Ala Ala Ala Ile Ile Gly Thr Asp Pro Ile Leu Asn Gln 225 230 235 240 Glu Ser Pro Phe Met Glu Leu Asn His Ala Val Gln Lys Phe Leu Pro 245 250 255 Asp Thr Gln Asn Val Ile Asp Gly Arg Ile Thr Glu Glu Gly Ile Asn 260 265 270 Phe Lys Leu Gly Arg Asp Leu Pro Gln Lys Ile Glu Asp Asn Ile Glu 275 280 285 Glu Phe Cys Lys Lys Ile Met Ala Lys Ser Asp Val Lys Glu Phe Asn 290 295 300 Asp Leu Phe Trp Ala Val His Pro Gly Gly Pro Ala Ile Leu Asn Lys 305 310 315 320 Leu Glu Asn Ile Leu Lys Leu Lys Ser Asp Lys Leu Asp Cys Ser Arg 325 330 335Lys Ala Leu Met Asp Tyr Gly Asn Val Ser Ser Asn Thr Ile Phe Tyr 340 345 350Val Met Glu Tyr Met Arg Asp Tyr Leu Lys Glu Asp Gly Ser Glu Glu 355 360 365 Trp Gly Leu Gly Leu Ala Phe Gly Pro Gly Ile Thr Phe Glu Gly Val 370 380

Leu Leu Arg Ser Leu

### M80676490.ST25

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270

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Gly Ile Ile Ser Lys Asn Ile Glu Lys Ser Leu Val Glu Ala Phe Ala 275 280 285

Pro Ile Gly Ile Asn Asp Trp Asn Ser Ile Phe Trp Val Ala His Pro 290 295 300

Gly Gly Pro Ala Ile Leu Asp Gln Val Glu Glu Lys Leu His Leu Lys 305 310 315 320

Glu Glu Lys Leu Arg Ser Thr Arg His Val Leu Ser Glu Tyr Gly Asn 325 330 335

Met Ser Ser Ala Cys Val Leu Phe Ile Leu Asp Glu Met Arg Lys Arg 340 345 350

Ser Lys Glu Glu Gly Met Ile Thr Thr Gly Glu Gly Leu Glu Trp Gly 355 360 365

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His Ser Val Pro Val Gln Gly 385 390

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#### M80676490.ST25

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Gly Tyr Ala Val Asn Thr Thr Val Arg Asp Pro Asp Ser Pro Lys Lys 35 40 45

Ile Ser His Leu Val Ala Leu Gln Ser Leu Gly Glu Leu Asn Leu Phe 50 60

Arg Ala Asp Leu Thr Val Glu Glu Asp Phe Asp Ala Pro Ile Ala Gly 65 70 75 80

Cys Glu Leu Val Phe Gln Leu Ala Thr Pro Val Asn Phe Ala Ser Gln 85 90 95

Asp Pro Glu Asn Asp Met Ile Lys Pro Ala Ile Lys Gly Val Leu Asn 100 105 110

Val Leu Lys Ala Ile Ala Arg Ala Lys Glu Val Lys Arg Val Ile Leu 115 120 125

Thr Ser Ser Ala Ala Ala Val Thr Ile Asn Glu Leu Lys Gly Thr Gly 130 140

His Val Met Asp Glu Thr Asn Trp Ser Asp Val Glu Phe Leu Asn Thr 145 150 155 160

Ala Lys Pro Pro Thr Trp Gly Tyr Pro Ala Ser Lys Met Leu Ala Glu 165 · 170 175

#### M80676490.ST25

Lys	Ala	Ala	Trp 180	Lys	Phe	ΑΊа	Glu	Glu 185	Asn	Asp	Ile	Asp	Leu 190	Ile	Thr
val	Ile	Pro 195	Ser	Leu	Thr	Thr	G]y 200	Pro	Ser	Leu	Thr	Pro 205	Asp	Ile	Pro
ser	ser 210	٧a٦	Gly	Leu.	Аla	Met 215	Ser	Leu	Ile	Thr	G]y 220	Asn	Asp	Phe	Leu
Ile 225	Asn	Ala	Leu	Lys	G]y 230	Met	Gln	Phe	Leu	ser 235	Glу	Ser	Leu	Ser	Ile 240
Thr	His	٧a٦	Glu	Asp 245	Ile	Cys	Arg	Ala	ніs 250	Ile	Phe	Leu	Ala	G]u 255	Lys
Glu	ser	Ala	ser 260	Gly	Arg.	Tyr	Ile	Cys 265	Cys	Ala	His	Asn	Thr 270	ser	∨al
Pro	Glu	Leu. 275	Ala	Lys	Phe	Leu	Asn 280	Lys	Arg	Tyr	Pro	G]n 285	Tyr	Lys	val
Pro	Thr 290	Glu	Phe	Asp	Asp	Cys 295	Pro	ser	Lys	Ala	Lys 300	Leu	Ile	Ile	Ser
ser 305	Glu	Lys	Leu	Ile	Lys 310	Glu	Gly	Phe	Ser	Phe 315	Lys.	His	Gly	Ile	Ala 320
Glu	Thr	Phe	Asp	G]n 325	Thr	Val	Glu	Tyr	Phe 330	Lys	Thr	Lys	Gly	Ala 335	Leu

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#### M80676490.ST25

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PRT.

Trifolium repens <213>

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Gly Arg Val Leu Ile Val Gly Gly Thr Gly Phe Ile Gly Lys Phe Val 20 25 30

Thr Glu Ala Ser Leu Ser Thr Thr His Pro Thr Tyr Leu Leu Val Arg 35 40 45

Pro Gly Pro Leu Leu Ser Ser Lys Ala Ala Thr Ile Lys Ala Phe Gln 50 60

Glu Lys Gly Ala Ile Val Ile Tyr Gly Arg Val Asn Asn Lys Glu Phe 65 70 75 80

Met Glu Met Ile Leu Lys Lys Tyr Glu Ile Asn Val Val Ile Ser Ala  $85 \hspace{1.5cm} 90 \hspace{1.5cm} 95$ 

Ile Gly Gly Ser Asp Gly Leu Leu Glu Gln Leu Thr Leu Val Glu Ala 100 105 110

Met Lys Ser Ile Asn Thr Ile Lys Arg Phe Leu Pro Ser Glu Phe Gly Page 14

M80676490.ST25 115 120 125

His Asp Val Asp Arg Ala Asn Pro Val Glu Pro Gly Leu Thr Met Tyr 130 140 Lys Gln Lys Arg Leu Val Arg Arg Val Ile Glu Glu Ser Gly Ile Pro 145 150 155 160 Tyr Thr Tyr Ile Cys Cys Asn Ser Ile Ala Ser Trp Pro Tyr Tyr Asp 165 170 175 Asn Cys His Pro Ser Gln Leu Pro Pro Pro Leu Asp Gln Leu His Ile 180 185 190 Tyr Gly His Gly Asp Val Lys Ala Tyr Phe Val Asp Gly Tyr Asp Ile 195 200 205 Gly Lys Phe Thr Met Lys Val Ile Asp Asp Glu Arg Thr Ile Asn Lys 210 220 Asn Val His Phe Arg Pro Ser Asn Asn Cys Tyr Ser Met Asn Glu Leu 225 230 235 240 Ala Ser Leu Trp Glu Asn Lys Ile Ala Arg Lys Ile Pro Arg Val Ile 245 250 255 Val Ser Glu Asp Asp Leu Leu Ala Ile Ala Ala Glu Asn Cys Ile Pro 260 265 270 Glu Ser Val Val Ala Pro Ile Thr His Asp Ile Phe Ile Asn Gly Cys 275 280 285 Gln Val Asn Phe Lys Ile Asp Gly Ile His Asp Val Glu Ile Gly Thr 290 295 300 Leu Tyr Pro Gly Glu Ser Val Arg Ser Leu Glu Glu Cys Tyr Glu Lys 305 310 315 320 Phe Val Val Met Ala Ala Asp Lys Ile His Lys Glu Glu Thr Gly Val Thr Ala Gly Gly Gly Thr Thr Ala Met Val Glu Pro Val Pro Ile 340 345 350 Thr Ala Ser Cys 355

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<sup>&</sup>lt;213> Trifolium repens

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1380	tttcatatat	tgtgttggtt	ttattgtagt	gtcttcatgt	tggggtaaaa	tattctctat
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1551	С	ctagtgaatt	tgcttaatca	ttgttaccac	gtactctgcg	aaaaaaaaa

Met Ala Pro Ala Ala Thr Ser Ser Pro Thr Thr Pro Thr Thr Lys  $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$ 

Gly Arg Val Leu Ile Val Gly Gly Thr Gly Phe Ile Gly Lys Phe Val  $20 \ \ 25 \ \ 30$ 

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#### M80676490.ST25

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45 Pro Gly Pro Leu Leu Ser Ser Lys Ala Ala Thr Ile Lys Ala Phe Gln 50 60 Glu Lys Gly Ala Ile Val Ile Tyr Gly Arg Val Asn Asn Lys Glu Phe 65 70 75. 80 Met Glu Met Ile Leu Lys Lys Tyr Glu Ile Asn Val Val Ile Ser Ala 85 90 95 Ile Gly Gly Ser Asp Gly Leu Leu Glu Gln Leu Thr Leu Val Glu Ala 100 105 Met Lys Ser Ile Asn Thr Ile Lys Arg Phe Leu Pro Ser Glu Phe Gly 115 120 His Asp Val Asp Arg Ala Asn Pro Val Glu Pro Gly Leu Thr Met Tyr 130 135 140 Lys Gln Lys Arg Leu Val Arg Arg Val Ile Glu Glu Ser Gly Val Pro 145 150 155 160 Tyr Thr Tyr Ile Cys Cys Asn Ser Ile Ala Ser Trp Pro Tyr Tyr Asp 165 170 175 Asn Cys His Pro Ser Gln Leu Pro Pro Pro Leu Asp Gln Leu His Ile 180 185 190 Tyr Gly His Gly Asp Val Lys Ala Tyr Phe Val Asp Gly Tyr Asp Ile 195 200 205 Gly Lys Phe Thr Met Lys Val Ile Asp Asp Glu Arg Thr Ile Asn Lys 210 220 Asn Val His Phe Arg Pro Ser Asn Asn Cys Tyr Ser Met Asn Glu Leu 225 230 235 240 Ala Ser Leu Trp Glu Asn Lys Ile Ala Arg Lys Ile Pro Arg Val Ile 245 250 255 Val Ser Glu Asp Asp Leu Leu Ala Ile Ala Ala Glu Asn Cys Ile Pro 260 265 270 Glu Ser Val Val Ala Ser Ile Thr His Asp Ile Phe Ile Asn Gly Cys 275 280 285 Gln Val Asn Phe Lys Val Asp Gly Ile His Asp Val Glu Ile Gly Thr 290 295 300

#### M80676490.ST25

Leu Tyr Pro Gly Glu Ser Val Arg Ser Leu Glu Glu Cys Tyr Glu Lys 305 310 315

Phe Val Val Met Ala Ala Asp Lys Ile His Lys Glu Glu Thr Gly Val

Thr Ala Gly Gly Gly Thr Thr Ala Met Val Glu Pro Val Pro Ile 340 345 350

Thr Ala Ser Cys 355

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#### M80676490.ST25

attc 1384

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<212> PRT <213> Trifolium repens

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Thr Glu Ala Ser Leu Ser Thr Thr His Pro Thr Tyr Leu Leu Val Arg 35. 40 45

Pro Gly Pro Leu Leu Ser Ser Lys Ala Ala Thr Ile Lys Ala Phe Gln 50 60

Glu Lys Gly Ala Ile Val Ile Tyr Gly Arg Val Asn Asn Lys Glu Phe 65 70 75 80

Met Glu Met Ile Leu Lys Lys Tyr Glu Ile Asn Val Val Ile Ser Ala 85 90 95

Ile Gly Gly Ser Asp Gly Leu Leu Glu Gln Leu Thr Leu Val Glu Ala 100 105 110

Met Lys Ser Ile Asn Thr Ile Lys Arg Phe Leu Pro Ser Glu Phe Gly 115 120

His Asp Val Asp Arg Ala Asp Pro Val Glu Pro Gly Leu Thr Met Tyr 130 140

Lys Gln Lys Arg Leu Val Arg Arg Val Ile Glu Glu Ser Gly Ile Pro 145 150 160

Tyr Thr Tyr Ile Cys Cys Asn Ser Ile Ala Ser Trp Pro Tyr Tyr Asp 165 170 175

Asn Cys His Pro Ser Gln Leu Pro Pro Pro Leu Asp Gln Leu His Ile 180 185 190

Tyr Gly His Gly Asp Val Lys Ala Tyr Phe Val Asp Gly Tyr Asp Ile 195 200 205

Gly Lys Phe Thr Met Lys Val Ile Asp Asp Glu Arg Thr Ile Asn Lys 210 220

Asn Val His Phe Arg Pro Ser Asn Asn Cys Tyr Ser Met Asn Glu Leu Page 19

> M80676490.ST25 235

230 225

240

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W	O 2004/090136		PCT/AU2004/000494
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			PCT/AU2004/	000494
Α.	CLASSIFICATION OF SUBJECT MATTER			
Int. Cl. <sup>7</sup> :	C12N 15/29	•		
According to I	nternational Patent Classification (IPC) or to both na	ational classification and IPC		
В.	FIELDS SEARCHED	,		
SEE ELECT	mentation searched (classification system followed by class			
SEE ELECT	searched other than minimum documentation to the exten RONIC DATABASES BELOW			ed
DGENE, Ger	base consulted during the international search (name of de nebank, EMBL, SwissProt, PIR, PubMed : SEC coanthocyanidin reductase, lcr, ban, dihydrofla	Q ID 2, 4, 6, 8, 10, 12, 14, 16	; Keywords: ch	alcone
C.	DOCUMENTS CONSIDERED TO BE RELEVANT			_
Category*	Citation of document, with indication, where appro	opriate, of the relevant passages		Relevant to claim No.
Р, Х	WO, A, 2003031622 (AGRICULTURE VICTAPRIL 2003 (SEQ ID 2 has 100% identity with with Fig. 148, SEQ ID 6 has 93% identity with Figure 163, SEQ ID 10 has 100% identity	n Figure 143, SEQ ID 4 has 9 th Figure 158, SEQ ID 8 has	6% identity	1-31
X WO, A, 2002057418 (THE SALK INSTITUTE FOR BIOLOGICAL STUDIES) 25 July 2002 (SEQ ID 2 has 92% identity with SEQ ID 1 on page 13)			1-3,6-8,10- 17,19-23,25- 29	
X SWISS-PROT database Accession Number P51088, Chalcone synthase 6. Howles, P. A. et al. 1 October 1996 (98% identity with SEQ ID 2)				1-3,6-8,10- 17,19-23,25- 29
X	SWISS-PROT database Accession Number P al. 1 October 1996 (97% identity with SEQ II	251083, Chalcone synthase 1. D 2)	Arioli, T., et.	1-3,6-8,10- 17,19-23,25- 29
X F	l urther documents are listed in the continuation	of Box C X See pa	tent family anne	×x
"A" documer not cons	idered to be of particular relevance cor unc	er document published after the international file with the application but cited to underlying the invention cument of particular relevance; the claim cannot be considered to involve an inv	nderstand the principion of the principion of the contract of	e or theory be considered novel
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)  "L" document which may throw doubts on priority claim(s) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art				be considered to one or more other
or other	nt published prior to the international filing date	cument member of the same patent fam	ily	
Date of the actu	than the priority date claimed nal completion of the international search	Date of mailing of the internation	al search report	JUL 2004
28 June 200		Authorized officer		
AUSTRALIAN	ing address of the ISA/AU I PATENT OFFICE	Authorized officer		
E-mail address	WODEN ACT 2606, AUSTRALIA : pot@ipaustralia.gov.au (02) 6285 3929	ALISTAIR BESTOW Telephone No: (02) 6283 245	,	
racsimile No.	(02) 0203 3727	reichione 140 . (02) 0205 245		

International application No.
PCT/AU2004/000494

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to
alegory	Citation of document, with indication, where appropriate, of the reaction process.	claim No.
X	GenPept database Accession Number CAA10131 Chalcone synthase, Dopico, B., et. al. 11 November 1998 (97% identity with SEQ ID 2)	1-3,6-8,10- 17,19 <b>-</b> 23,25 29
X	SWISS-PROT database Accession Number Q9SEVO Leucoanthocyanidin reductase, Devic, M., et. al. 28 February 2003 (61% identity with SEQ ID 10)	1,2,4,6,7,9- 16,18-22,24 28,30
X	GenPept database Accession Number AAF23859 11 January 2000 (61% identity with SEQ ID 10)	1,2,4,6,7,9- 16,18-22,24 28,30
X	JEZ, J. M. et.al. Structure and Mechanism of Chalcone Synthase-like Polyketide Synthases, <i>Journal of Industrial Microbiology &amp; Biotechnology</i> (2001) 27:393-8	1-3,6-8,10- 17,19-23,25 29
X	ARIOLI, T., et.al. In <i>Trifolium subterraneum</i> , chalcone synthase is encoded by a multigene family. Gene, 138:79-86 (1994) (SEQ ID 2 is 97% identity to seq. of Fig. 3)	1-3,6-8,10- 17,19-23,25 29
X	CHARRIER, B., et.al. Molecular characterization and expression of alfalfa ( <i>Medicago sativa</i> L.) flavanone-3-hydroxylase and dihydroflavonol-4-reductase encoding genes. Plant Molecular Biology 1995 Nov;29(4):773-86 (See Figure 3)	1,2,4,6,7,9- 16,18-22,24 28,30
X	WO A1 2002066625 (COMMONWEALTH SCIENTIFIC AND INDUSTRIAL RESEARCH ORGANISATION ET. AL.) 29 August 2002 (SEQ ID 12 and 14 and 16 have 72% identity to SEQ ID 28)	1,2,5,6,8- 15,17-21,23 28,31.
X	WO A1 2000022099 (GENESIS RESEARCH AND DEVELOPMENT CORPORATION LIMITED et. al.) 20 April 2000, (SEQ ID 12 and 14 have 56% identity to SEQ ID 323. SEQ ID 16 has 57% identity to SEQ ID 323)	1,2,5,6,8- 15,17-21,23 28,31
X	EP A 1033405 (CERES INC.) 6 September 2000. (SEQ ID 4 has 77% identity with SEQ ID 66257 and SEQ ID 15839, SEQ ID 6 has 67% identity with SEQ ID 7420 and 7419)	1-3,6-8,10- 17,19-23,25 29
X	WO A 2002010210 (BAYER AG) 7 February 2002. (SEQ ID 4 has 77% identity with SEQ ID 2451, SEQ ID 6 has 70% identity with SEQ ID 18.)	1-3,6-8,10- 17,19-23,25 29
X	EP A 1254960 (UNILEVER PLC) 6 November 2002. (SEQ ID 8 has 81% identity with Fig.11.)	1-3,6-8,10- 17,19-23,25 29
P,X	WO A 2003040306 (GENESIS RESEARCH AND & DEVELOPMENT CORPORATION LTD) 15 May 2003. (SEQ ID 8 has 78% identity with SEQ ID 119 and SEQ ID 186.)	1-3,6-8,10- 17,19-23,25 29
P,X	WO A 2004020637 (INTERNATIONAL FLOWER DEVELOPMENTS PTY. LTD.) 19 March 2004. (SEQ ID 8 has 80% identity with disclosure on p.244-6.)	1-3,6-8,10- 17,19-23,25 29
Χ.	PIR database Accession Number T10231 Anther-specific protein homolog T11I11.90, Bevan, M., et. al. 16 July 1999. (77% identity with SEQ ID 4.)	1-3,6-8,10- 17,19-23,25 29

International application No.

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X .	PIR database Accession Number T15054. Anther-specific protein - wood tobacco. Atanassov, I.I., et. al. 20 September 1999. (76% identity with SEQ ID 6.)	1-3,6-8,10- 17,19-23,25 29
Χ.	EMBL database Accession Number Y14507, Nicotiana sylvestris CHSLK gene. Atanassov I.I., et. al. 11 August 1997. (70% identity with SEQ ID 6.)	1-3,6-8,10- 17,19-23,25 29
X	SWISS-PROT database Accession Number P51075, Chalcone synthase. Pellinen, R., et. al. 1 October 1996. (82% identity with SEQ ID 8.)	1-3,6-8,10- 17,19-23,25 29
х	SWISS-PROT database Accession Number Q9FSB7, Chalcone synthase 3. Springob, K., et. al. (82% identity with SEQ ID 8), and, SPRINGOB, K., et. al. Specificities of functionally expressed chalcone and acridone synthases from <i>Ruta graveolens</i> , Eur. J. Biochem. 267(22):6552-9 (2000)	1-3,6-8,10- 17,19-23,25 29
X	TREMBL database Accesssion Number BAB84112. Chalcone synthase (Vitis vinifera). Goto-Yamamoto, N., et. al. 19 July 2002. (82% identity with SEQ ID 8.)	1-3,6-8,10- 17,19-23,25 29
		·

International application No.

Во	x No.	Ι	Nucleotide and/or amino acid sequence(s) (Continuation of item 1.b of the first sheet)
1.	With	n rega ned ir	rd to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the evention, the international search was carried out on the basis of:
	a.	type	of material
		X	a sequence listing
			table(s) related to the sequence listing
	ъ.	forma	at of material
			in written format
	•	X	in computer readable form
	c.	time	of filing/furnishing
		X	contained in the international application as filed
			filed together with the international application in computer readable form
			furnished subsequently to this Authority for the purposes of search
2.		file	ddition, in the case that more than one version or copy of a sequence listing and/or table(s) relating thereto has been d or furnished, the required statements that the information in the subsequent or additional copies is identical to that in application as filed or does not go beyond the application as filed, as appropriate, were furnished.
3.	Addi	tional	comments:
,			

International application No.

Box No	. П	Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This interessons:		ational search report has not been established in respect of certain claims under Article 17(2)(a) for the following
1.		Claims Nos.:
		because they relate to subject matter not required to be searched by this Authority, namely:
2.	$\Box$	Claims Nos.:
_	-	because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.		Claims Nos.:
		because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)
Box No	. m	Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
This Int	ern	ational Searching Authority found multiple inventions in this international application, as follows:
Applica inventi	atic	cant has claimed more than one invention. Rule 13.1 of the PCT states the principle that an International on should relate to only one invention or, if there is more than one invention, that the inclusion of those in one International Application is only permitted if all inventions are so linked as to form a single general concept.
•		(continued on Extra Sheet)
1. 2	ζ]	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.		As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.		As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4.		No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remarl	k or	The additional search fees were accompanied by the applicant's protest.
		X No protest accompanied the payment of additional search fees.

International application No.

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Supplemental Box

(To be used when the space in any of Boxes I to VIII is not sufficient)

#### Continuation of Box No: III

Rule 13.2 of the PCT defines the method for determining whether the requirement of unity of invention is satisfied in respect of a group of inventions claimed in an International application. Unity of invention exists only when there is a technical relationship among the claimed inventions involving one or more of the same or corresponding "special technical features." The expression "special technical features" is defined in Rule 13.2 as meaning those technical features that define a contribution which each of the inventions, considered as a whole, makes over the prior art. The determination is made on the contents of the claims as interpreted in light of the description and drawings (if any).

**Invention 1**: Polypeptide and polynucleotide encoding chalcone synthase of the formula TrCHSa3, represented by SEQ ID NOS 1 and 2. This invention is recited in claims 1 - 3, 6 - 8, 10 - 17, 19 - 23, 25 - 29 (in part).

Invention 2: Polypeptide and polynucleotide encoding chalcone synthase of the formula TrCHSc, represented by SEQ ID NOS 3 and 4. This invention is recited in claims 1 - 3, 6 - 8, 10 - 17, 19 - 23, 25 - 29 (in part).

Invention 3: Polypeptide and polynucleotide encoding chalcone synthase of the formula TrCHSf, represented by SEQ ID NOS 5 and 6. This invention is recited in claims 1 - 3, 6 - 8, 10 - 17, 19 - 23, 25 - 29 (in part).

Invention 4: Polypeptide and polynucleotide encoding chalcone synthase of the formula TrCHSh, represented by SEQ ID NOS 7 and 8. This invention is recited in claims 1 - 3, 6 - 8, 10 - 17, 19 - 23, 25 - 29 (in part).

Invention 5: Polypeptide and polynucleotide encoding dihydroflavonol 4-reductase of the formula TrBANa, represented by SEQ ID NOS 9 and 10. This invention is recited in claims 1, 2, 4, 6 - 7, 9, 10 - 16, 18 - 22, 24 - 30. (in part)

Invention 6: Polypeptide and polynucleotide encoding leucoanthrocyanine reductase of the formula TrLARa, represented by SEQ ID NOS 11 and 12. This invention is recited in claims 1, 2, 5, 6, 8 - 15, 17 - 21, 23 - 26, 28, 29, 30. (in part).

**Invention 7**: Polypeptide and polynucleotide encoding leucoanthrocyanine reductase of the formula TrLARb, represented by SEQ ID NOS 13 and 14. This invention is recited in claims 1, 2, 5, 6, 8 - 15, 17 - 21, 23 - 26, 28, 29, 30. (in part).

Invention 8: Polypeptide and polynucleotide encoding leucoanthrocyanine reductase of the formula TrLARc, represented by SEQ ID NOS 15 and 16. This invention is recited in claims 1, 2, 5, 6, 8 - 15, 17 - 21, 23 - 26, 28, 29, 30. (in part).

Each of the above enzymes may be involved in flavonoid biosynthesis in plants, and more specifically the modification of the content of condensed tannins, but this not novel, as noted in the following abstract. Therefore this cannot be used as a special technical feature providing unity to all of the sequences.

XIE DY et. al. Science (Jan 2003)299(5605):396-9 Role of anthocyanin reductase, encoded by BANYULS in plant flavonoid biosynthesis.

Although some of the claims share a technical feature whereby they are a chalcone synthase, a dihydroflavonol 4-reductase or a leucoanthocyanine reductase, these are not special technical features because none of these are novel enzymes. For example, the following three journal abstracts indicate that none of these enzymes are novel, and therefore cannot be used as a special technical feature uniting the inventions into three groups.

NAPOLI C, et. al. The Plant Cell (1990) 2:279-89 Introduction of a Chimeric Chalcone Synthase Gene into Petunia Results in Reversible Co-suppression of Homologous Genes *in trans*.

JOHNSON ET, et. al. Plant Journal (2001) 25(3):325-33 Alteration of a single amino acid changes the substrate specificity of dihydroflavonol 4-reductase.

TANNER GJ et.al. Anal Biochem 1993 March 209(2):274-7 Synthesis of 3,4-cis-[3H]leucocyanidin and enzymatic reduction to catechin

(Continued on Extra Sheet.)

International application No.

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Supp	lemen	tal	Box
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(To be used when the space in any of Boxes I to VIII is not sufficient)

#### Continuation of Box No:

Furthermore, given the nature of the claims and the invention, it is appropriate to apply the Markush approach. Using the Markush approach to analyse the unity of inventions, although the 8 gene transcripts all have the common property (A) that they are involved in flavonoid biosynthesis in plants, and more specifically the modification of the content of condensed tannins, (B)(1) there is no common structure present in all of the genes; and (B)(2) there is no single recognised class or group of compounds embracing all the genes claimed. It is contrary to normal classification to group together such diverse genes. Thus according to Markush, it is appropriate to classify the genes in terms of the 8 individual groups and thus these groups represent 8 different inventions.

recognised class or group of compounds embracing all the genes claimed. It is contrary to normal classification to group together such diverse genes. Thus according to Markush, it is appropriate to classify the genes in terms of the 8 individual groups and thus these groups represent 8 different inventions.							
In order to search each of the inventions, this could only be done by consideration of each of the sequences, thereby requiring eight separate searches. As a service to the applicant, this office would for the purposes of this search, consider Inventions 1 and 5 together for a single search fee. Each of the other sequences will be considered as single inventions and each attracts a search fee.							
The applicant chose to pay the six extra fees being sufficient to search and examine all eight inventions.							
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